EAST Search History

Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	0	("2004006104").PN.	US-PGPUB	OR	OFF	2006/05/11 07:14
L2	1	("20040006104").PN.	US-PGPUB	OR	OFF	2006/05/11 07:19
L3	1	("6482949").PN.	US-PGPUB; USPAT	OR	OFF	2006/05/11 08:30
L4	5	"2005007099"	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2006/05/11 08:30
S1	985	544/344 OR 544/347 OR 544/353	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	OFF	2006/05/11 07:13
S2	72	S1 AND (ANTIVIRAL OR HCV OR HEPATITIS)	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	ON	2006/05/10 11:58
S3	1	("6989451").PN.	USPAT	OR	OFF	2006/05/10 12:31
S4	1	("3510487").PN.	USPAT	OR	OFF	2006/05/10 12:42
S5	1	("3656953").PN.	USPAT	OR	OFF	2006/05/10 13:29
S6	1	("6518423").PN.	USPAT	OR	OFF	2006/05/10 13:39
S7	1	("6103720").PN.	USPAT	OR	OFF	2006/05/10 13:43
S8	1	("5874587").PN.	USPAT	OR	OFF	2006/05/10 13:45
S9	1	("5969150").PN.	USPAT	OR	OFF	2006/05/10 13:45

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property data NEWS 6 JAN 17 NEWS 7 JAN 17 NEWS 8 JAN 30 NEWS 9 FEB 21 NEWS 15 FEB 28 REDISTRY/ZREDISTRY enhanced with more experimental spectral NEWS 15 FEB 28 REDISTRY/ZREDISTRY enhanced with more experimental spectral NEWS 16 MAR 01 INSPEC reloaded and enhanced NEWS 17 MAR 03 Updates in PATDPA; addition of IPC 8 data without attributes NEWS 17 MAR 08 K.25 communication option no longer available after June 2006 NEWS 19 MAR 28 EMBASE is now updated on a daily basis NEWS 20 APR 03 New IPC 6 fields and IPC thesaurus added to PATDPAPULL NEWS 21 APR 03 Bibliographic data updates resume; new IPC 8 fields and IPC thesaurus added in PCTFULL NEWS 22 APR 03 STN Anavist \$500 visualization usage credit offered NEWS 23 APR 12 LINSPEC, learning database for INSPEC, reloaded and enhanced NEWS 24 APR 12 Improved structure highlighting in PQHIT and QHIT display in MQNAPT
NEWS 25 APR 12 Dervent World Patents Index to be reloaded and enhanced during second querter; strategies may be affected
NEWS 26 MAY 10 CA/CAplus enhanced with 1900-1906 U.S. patent records

NEMS EXPRESS FEBRUARY 15 CURRENT VERSION FOR WINDOWS IS V8.01a,
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chain nodes:
11 13 15 16 18 19
ring nodes:
1 2 3 4 5 6 7 8 9 10
ring/chain nodes:
20 chain bonds : 5-11 11-13 11-15 15-16 16-18 16-19 18-20 5-11 11-13 11-15 15-16 16-16 16-17 20-17 17-17 18-18 18-19 18-18 1

G2:0.N

G3:C.H.S.P

Match level : 1:Atom 3:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:CLASS 13:CLASS 15:CLASS 16:CLASS 18:CLASS 19:CLASS 20:CLASS

Ll STRUCTURE UPLOADED

-> que Ll

L2 QUE L1

-> D L1 L1 HAS NO ANSWERS

THIS IS FOR CL. 151, × 3, 4, 6

G1 C, S G2 O.N G3 [01], [02]

Structure attributes must be viewed using STN Express query preparation.

SAMPLE SEARCH INITIATED 12:13:34 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 530 TO ITERATE

100.0% PROCESSED 530 ITERATIONS SEARCH TIME: 00.00.01

12 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
PROJECTED ITERATIONS: 9219 TO 11981
PROJECTED ANSWERS: 33 TO 447

L3 12 SEA SSS SAM L1

-> S L1 SSS PULL FULL SEARCH INITIATED 12:13:40 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 10767 TO ITERATE

100.04 PROCESSED 10767 ITERATIONS SEARCH TIME: 00.00.01

329 ANSWERS

329 SEA SSS FUL L1

->Testing the current file.... screen

ENTER SCREEN EXPRESSION OR (END) : end

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chain nodes:
11 13 15 16 18 19
ring nodes:
1 2 3 4 5 6 7 8 9 10
ring/chain nodes:

G2:0.N G3:C,H,S,P

Match level: 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:CLASS 13:CLASS 15:CLASS 16:CLASS 16:CLASS 19:CLASS 20:CLASS

L5 STRUCTURE UPLOADED

-> que L5 L6 QUE L5

-> D L5 L5 HAS NO ANSWERS L5

Structure attributes must be viewed using STN Express query preparation.

SAMPLE SEARCH INITIATED 12:16:14 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 530 TO ITERATE

100.0% PROCESSED 530 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

50 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
PROJECTED ITERATIONS: 9219 TO 11981
PROJECTED ANSWERS: 933 TO 1607

50 SEA SSS SAM LS

G2 O, N G3 C, H, S, P

-> S L5 SSS PULL
FULL SEARCH INITIATED 12:16:19 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 10767 TO ITERATE

100.0% PROCESSED 10767 ITERATIONS SEARCH TIME: 00.00.01

1540 ANSWERS

1540 SEA SSS PUL L5

-> S L8 NOT L4 L9 1213 L8 NOT L4

->Testing the current file.... screen

ENTER SCREEN EXPRESSION OR (END) : end

Uploading C:\Program Files\Stnexp\Queries\ANRONGWUCHOW ANTIVIRALS.str

chain nodes:
11 13 15 16 16 19 ring nodes:
1 2 3 4 5 6 7 8 9 10 ring/chain nodes:
20 20 chain bonds :
5-11 11-13 11-15 15-16 16-18 16-19 18-20 ring bonds :
1-2 1-6 2-3 2-7 3-4 3-10 4-5 5-6 7-8 8-9 9-10 exact/norm bonds :
11-13 11-15 15-16 exact bonds :
5-11 16-18 16-19 18-20 normalized bonds :
1-2 1-6 2-3 2-7 3-4 3-10 4-5 5-6 7-8 8-9 9-10

G2:0, N

03 · C. H. S. P. Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:CLASS 13:CLASS 15:CLASS 16:CLASS 18:CLASS 19:CLASS 20:CLASS

L10 STRUCTURE UPLOADED

-> que L10

L11 QUE L10

a> D L10 L10 HAS NO ANSWERS L10 STR

G1 C,S G2 O, N

Structure attributes must be viewed using STN Express query preparation.

=> S L10 SUB-L9 FULL FULL SUBSET SEARCH INITIATED 12:21:05 FILE 'REGISTRY' FULL SUBSET SCREEN SEARCH COMPLETED - 987 TO ITERATE

SEARCH TIME: 00.00.01 987 ITERATIONS 790 ANSWERS 790 SEA SUB-L9 SSS FUL L10 L12 FULL ESTIMATED COST

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-> 8 L4 OR L12

151 L12 L1 OR L12 — ALL SEARCHES TOGETHER

DESPLAYED 1- 18/1.

L13 ANSWER 1 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2006;36:1356 CAPLUS
TITLB: Preparation of disulfide dyes for dyeing human keratin fibers

Daubresse, Nicolas: Genain, Gilles INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE: Fr. U.S. Pat. Appl. Publ., 39 pp. CODEN: USXXCO

Patent English DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO.

PATENT NO. KIND DATE APPLICATION NO. DATE

US 2006080791 A1 20060420 US 2005-249357 20051014

FR 2876576 A1 20060421 FR 2004-10864 20041014

EP 1647580 A1 20060419 EP 2005-293159 20051013

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BO, CZ, EE, RU, PL, SK,

BA, HR, IS, YU

JP 2006111626 A2 20060427 JP 2005-299150 20051013

ENTY APPLA INFO: FR 2004-10864 A 20041014

PRIORITY APPLN. INFO.:

JP 2006111626 A2 20060427 JP 2005-299150 20051013
RRITY APPLN. INFO.: PR 2004-10864 A 20041014
Disclosed herein is a dyeing composition comprising a particular disulfide dye and a method of dyeing human keratin fibers, such as hatr, using this composition this composition makes it possible to obtain particularly fast chromatic colorations. E.g., I was prepared from cyclamine-2HCl and Reactive Blue 44. I and other prepared dyes were tested on gray hair. INDEXIND IN PROGRESS 803566-67-4P
EL: SEU (Biological applied)

osjoob-e/-y RK: BSU (Biological study, unclassified); COS (Cosmetic use); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES

(Uses)
(preparation of disulfide dyes for dyeing human keratin fibers)
88356-67-4 CAPLUS
INDEX NAMEN FOT YET ASSIGNED

DBGG 1.A

This invention relates to novel compds, useful in the treatment of diseases associated with TRPV4 channel receptor. E.g., I was prepared from Z-D-DAB(BOC)-OH DCHA and ClCOZEt giving an intermediate which was treated with phthelimide and Ph3P and DBAD giving phenylaethyl (2R) 4-[[(1.1-dimethylethoxy)carbonyl]maino]-2-[(1.3-dioxo-1,3-dihydro-2H-isoindol-2-ylmethyl]butaneate which was treated with RCl and N-(1-benzothien-2-ylearbonyl)-L-leucine. The resulting intermediate was brominated and treated with 2-chloro-4-fluorobenzenesulfonyl chloride and the resulting intermediate hydrazinolyzed to give I. Tablets were prepared containing I. the resulting intermediate hydracardy and the containing I.
878797-73-09
RL: SPN (Symthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of acyclic 1, 3-diamines for use in treatment of diseases associated with TRPV4 channel receptor)

878797-73-0 CAPLUS
6-Quinoxalinecarboxamide, N-[(1S)-1-[[(3-[[(2-chloro-4-fluorophenyl)aulfonyl]amino]propyl)amino]carbonyl]-3-methylbutyl]- (9CI)
(CA INDEX NAME)

Absolute stereochemistry

PUBLISHER: DOCUMENT TYPE: LANGUAGE: GI

L13 ANSNER 3 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
TITLE:
Nucleic exid intercalators and avidin probes derived from luminescent cyclometalated iridium(III)dipyridoquinoxaline and -dipyridophenazine complexes
LO, Kenneth Kama-Wing; Chung, Chi-Keung; Zhu, Nianyong
Department of Biology and Chemistry, City University of Hong Kong, Kovloon, Hong Kong, Poop. Rep. China
Chemistry--A European Journal (2006), 12(5), 1500-1512
CODEN: CEUISD: ISSN: 0947-6539
UNILISHER:
MILey-VCH Verlag GmbH & Co. KGeA
Journal
GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

L13 ANSWER 2 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2006:240647 CAPLUS DOCUMENT NUMBER: 144:311390 Preparation of accession of accession and accession access 144:311900

Preparation of acyclic 1,3-diamines for use in treatment of diseases associated with TRPV4 channel receptor.

Casillas, Linda N.; Jeong, Jae UK; Marquis, Robert W. Smithkline Beecham Corporation, USA PCT Int. Appl., 152 pp.
CODEN: PIXXD2
Patent

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: English 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

MO 2006029210 A2 20060316 M0 2005-US31873 20050997
M: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CM, CM, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, EG, ET, GB, GD, GB, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KP, KR, KZ, LC, LK, LR, LS, IT, LU, LV, MA, MD, MG, KK, MN, MM, MX, AZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RG, RU, SC, SD, SE, SG, EK, SL, SM, SY, TJ, TM, TM, TR, TT, TZ, LA, UG, US, UZ, VC, VN, VU, ZA, ZM, ZM, ZM
RM: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NI, PL, PT, RG, SE, SI, SK, TR, BF, BJ, CP, CG, CI, CM, GA, GM, GQ, GM, ML, MR, NE, SM, SN, TD, TG, BM, GH, GM, KE, LS, MM, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZM, AM, AZ, BY, KUT APPLIN. INFO: KG, KZ, M PRIORITY APPLN. INFO.: GI

Six luminescent cyclometalated cationic redox-active luminescent
2-phenylpyridine iridium(III)-complexes with substituted
pyrazino-annelated phenanthroline bidentate ligands were prepared; the DNA
and avidin intercalation were assayed by emission titration Reaction of
[Ir2(ppy)4Cl2] (ppy = 2-phenylpyridine) with ligands L2 gave complexes
1-P96 [1-3; L2 = dpq R = H; L2 = dpqa, R = CONNHSU, L2 = dpqS, R =
CONN(CH2)2NMO, where O = CO(CH2)4CSHYNOS, biotinyl] and II [4-6; L2 =
dpps, R1 = R2 = H; L2 = dppn, R1-R2 = benco, L2 = dpp3, R2 = R1 =
CONN(CH2)2NMO) were designed as luminescent intercalators for DNA and
probes for avidin. The crystal structure of complex 4 is reported. The
photophys. and electrochem. properties of the complexes 1-6 were also
investigated. The binding of these complexes to double-stranded calf
thymus DNA and synthetic double-stranded oligonucleotides
poly(AL)-poly(dT) and poly(dG)-poly(dC) was investigated by
spectroscopic titrms. The interactions between the two biotin-containing
complexes 3 and 6 and avidin were studied by 4-hydroxysobenzene-2complexes 3 and 6 and avidin were studied by 4-hydroxysobenzene-2septimal complexants (RNAEM) assays and emission titrms.
SE2571-97-30
SEL: RCT (Reactant): SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or resgent)
(preparation of cyclometalated luminescent iridium pyrszino-annelated
phenanthroline 2-phenylpyridine complexes as DNA intercalstors and
avidin complexants
SE2571-97-3 CAPLUS
Ouinoxalino[2,3-f][1,10]phenanthroline-11-carboxamide,
N-[2-[5-[3as,48,6ar]-hexhydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl]-1oxopentyllamino[ethyl] (SCI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

882571-95-1P
RL: 88U (Biological study, unclsseified); CPS (Chemical process); PEP
(Physical, engineering or chemical process); PTP (Physical process); SPN
(Synthetic preparation); BIOL (Biological study); PRSF (Preparation); PROC
(Process)

Occass)
{redox potential, luminescence spectrs; preparation of cyclometalated
luminescent iridium pyraxino-annelated phenanthroline 2-phenylpyridine

complexes as DNA intercalators and avidin complexants) 862571-95-1 CAPLUS INDEX NAME NOT YET ASSIGNED

CM 1

CRN 882571-94-0 CMF C53 H46 Ir N10 O3 S CCI CCS

PAGE 1-A

A 20040809 A 20040927 A 20050322 A 20030930 A 20040310 WO 2004-JP14063 JP 2005-82760 OTHER SOURCE(s):

MARPAT 144:232928

Antimalaria agents containing compds. represented by the formula (I) (wherein Al = each optionally substituted 3-pyridyl or 6-quinolyl; Xl = -C('Xl)-NN-; Yl = 0; B = each optionally substituted furyl, thienyl, or phenyl; provided that Al may have one to three substituents and B has one or two substituents), salts of the compds., or hydrates of either are disclosed. Thus, a solution of 2-mainonicotinic acid and [[5-(3-chlorobenzyl)furan-2-yl]methyl]mains in DNF was treated with benzotriazol-1-yl-trie (dimethylamino) phosphonium hazaflorosphosphate and EtN and stirred at 80° for 40 min to give 3-maino-N-[5-(3-chlorobenzyl)furan-2-ylenthyl]neotimanide [II]. II showed min. inhibitory concentration of 6.25 µg/mL against yeast expressing plasmodium GNT1 gene (pp(GNT1).
849810-87-1P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapoutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of heterocyclic compds. such as nicotinamide quinolinecarboxamide derive. as antimalaria agents)
849810-87-3 CAPUUS
6-Quinoxalinecarboxamide N-((3-phenexyphenyl)methyl)-

849810-87-3 CAPLUS
6-Quinoxalinecarboxamide, N-[(3-phenoxyphenyl)methyl]-,
mono(trifluoroacetate) (9CI) (CA INDEX NAME)

CM 2 16919-18-9 P6 P CCS

REFERENCE COUNT:

THERE ARE 115 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT

L13 ANSWER 4 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2006:152549 CAPLUS
DICTURENT NUMBER: 144:23228
TITLE: PATENT ASSIGNEE(S): Responsible to the compound of the compound o

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

CRN 849810-86-2 CMF C22 H17 N3 O2

CRN 76-05-1 CMF C2 H F3 O2

c-co₂н

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

English

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

MO 2006005911 Al 20060119 MO 2005-082729 200507311

M: AS, AO, AL, AM, AT, AU, AZ, AD, BB, BO, BR, BM, BY, BZ, CA, CH, CO, CC, CR, CU, CZ, DR, DK, DM, DZ, EC, ER, EO, ES, FI, GB, GD, GB, GM, HR, HU, JD, IL, IN, 1S, JP, KE, KG, CM, KP, KR, KZ, CM, KD, CL, LK, LR, LS, LT, LU, LV, MA, MD, MG, MN, MM, MM, MX, MZ, NA, MO, NI, NO, NZ, OM, PQ, PH, PL, PT, RO, RU, SC, SD, ES, SO, GK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UO, US, UZ, VC, VN, VU, ZA, ZM, ZM

RM: AT, BR, BG, CH, CY, CZ, DE, DK, ES, FI, FR, GB, GR, HU, IE, IE, IT, LT, LU, LV, MC, NL, PL, PT, RO, RS, SI, SK, TR, BF, BJ, CP, CO, CI, CM, GA, GN, GQ, GM, ML, MR, NS, SN, TD, TO, BM, GH, GM, KE, LS, MM, MZ, NA, SD, SL, ZT, ZT, UG, XL, ZM, AM, AZ, BY. PRIORITY APPLAN. INPO:: US 2004-587177P

OTHER SOURCE(S):

MARPAT 144:150635

B The invention relates to compds. R1(CH2)0-3NR5COCH[NR4-X-(CH2)0-3R3] (CH2)3-6COR2 [X is CH2, CO, SO2, CONH, CO2, C(S)NH OT CONMSO2; R1 is (un)substituted carbalkoxy, amino groups, aryl, aryloxy, cycloskyl, aryl or heterocyclyl; R2 is H, (un)substituted alkyl, carbamoyl, CF3, cycloskyl, aryl or heterocyclyl; R3 is H, CF3, Oxo, OH, CN, halo, amino groups, (un)substituted carboxylic ester, acyl, sulfonyl groups, etc.; R4 is H or alkyl; R5 is H or together with R1(CH2)0-3N forms (un)substituted piperaxinyl] that are inhibitors of histone deacetylase (HDAC) and are useful for treating cancer, neurodegenerative diseases, schizophrenia, atroke and other diseases. Thus, (2S)-2-[(5-methoxy-2-methyl-1H-indol-3-yl)actyl]smino-8-oxo-N-[2-12-phenyl-1H-indol-3-yl)-lethyl]nonamide was prepared by a multistep sequence involving reactions of Me 8-oxonomanoate, ethylendeiol, (S)-()-4-benyl-2-oxazolidinone, 2-(2-phenyl-1H-indol-3-yl)-thanaminius chloride, and 5-methoxy-2-methyl-3-indolylacetic acid. Compds. of the invention were found to have HDAC inhibitory activity (ICS0 of 10 MM).

18 27454-44-69 874159-11-2P 874159-77-0P 874159-15-19 874159-17-9P 874159-77-0P 874159-17-9P 874159-20-20-3P ML: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); Objective of histone

(Uses)
(preparation of amino acid amide derive. as inhibitors of histone deacetylase)
deacetylase)
6-QUINOS
6-QUINOSALInecarboxamide, N-[(18)-7-oxo-1-[[[2-(2-phenyl-1H-indol-3-yl)ethyl]amino]carbonyl]octyl]- (9CI) (CA INDEX NAME)

Absolute atereochemiatry.

874159-11-2 CAPLUS 6-Quinoxalinecarboxamide, N-[(1S)-1-[[(3-acetylphenyl)amino]carbonyl]-7-oxononyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

874159-77-0 CAPLUS 6-Quinoxalinecarboxamide, N-[(18)-1-[(2-naphthalenylamino)carbonyl]-7-oxononyl]- (9CI) (CA INDEX NAME)

874160-17-5 CAPLUS 6-Quinoxalinecarboxamide, N-[(1S)-7-oxo-1-[[(2-phenylethyl)amino]carbonyl]nonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

874160-23-3 CAPLUS 6-Quinoxalinecarboxamide, N-[(18)-7-oxo-1-[[[2-(3-phenyl-1-pyrrolidinyl)ethyl]amino]carbonyl]nonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

874159-15-6 CAPLUS 6-Quinoxalinecarboxamide, N-[(18)-1-[(cyclopentylamino)carbonyl]-7-oxnonyl]- (SCI) (CA INDEX NAME)

Absolute stereochemistry.

874159-36-1 CAPLUS 6-Quinoxalinecarboxamide, N-[(18)-7-oxo-1-[(3-pyridinylamino)carbonyl]nonyl]- (9CI) (CA INDEX NAME)

874159-71-4 CAPLUS 6-Quinoxalinecarboxamide, N-[(1S)-7-oxo-1-[[[(2-phenyl-4-thiazolyl)methyl]amino]carbonyl]nonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 6 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2006:16547 CAPLUS

DOCUMENT NUMBER:

CORPORATE SOURCE:

SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

OTHER SOURCE(S):

MONET TYPE: Journal
NUMBE: Elsevier B.V.
MENT TYPE: Journal
NUMBE: Anglish
RE SOURCE(S): CASREACT 144:274236
A set of 29 6-aminoquinoxalines and 6-quinoxalinecarboxamides are prepared
and evaluated in vitro against several protozoal paraeites (Leishmania
donovani, Trypanoxome brucei brucei, and Trichomonas vaginalis); four
compds. are active as antileishmanial agents with ICSO values of < 20
µM. While none of the brominated quinoxalines or 2,3diphenylquinoxalines prepared are active as antiprotozoal agents, no other
clear structure-activity relationahip among the quinoxalinea prepared ia
found.

found.
979330-07-4P 979230-12-1P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL
(Biological atudy); PREP (Preparation)
(preparation of aminoquinoxalines and quinoxalinecarboxamidea, their
antiprotozeal atructure-activity relationshipa, and their activities
against Leishmania donovani, Trypanoaoma brucei brucei, and Trichomonae
vaginalia)
878230-07-4 CAPLUS
6-Quinoxalinecarboxamide, N-[(2,3-diphenyl-6-quinoxalinyl)carbonyl]-N-(3methoxy-2-dibenzofuranyl)-3,3-diphenyl- (9CI) (CA INDEX NAME)

878290-12-1 CAPLUS 6-Quinoxalinecarboxamide, N-[(2,3-diphenyl-6-quinoxalinyl)carbonyl]-N-(9-ethyl-9H-carbazol-3-yl)-2,3-diphenyl- (9CI) (CA INDEX NAME)

THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 7 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2005:1350084 CAPLUS

DOCUMENT NUMBER: TITLE:

2005:1350084 CAPLUS
144:88701
Charge-transport materials, methods of fabrication thereof, and methods of use thereof
Marder, Seth; Kaafarani, Bilal; Barlow, Stave;
Kippelen, Bernhard; Domercq, Benoit; Zhang, Qing;
Kondo, Taksehi
Georgia Fech Research Corporation, USA
PCT Int. Appl., 219 pp.
CODEN: PIXXD2
Patent
1

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. KIND A2 C2 PATENT NO. DATE WO 2005123737 WO 2005123737 20051229 WO 2005-US20998 20050614 2005123737 C2 20050006
M: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BM, BY, BZ, CA, CH, CM, CO, CR, CU, CZ, DR, DK, DM, DZ, EC, RE, EG, ES, FT, GB, GD, GB, GH, GH, HB, HU, 1D, IL, IN, 1S, JP, ER, KG, KM, KP, KR, KZ, LC, LK, LR, LB, LT, LU, LV, NA, MD, MG, MK, MN, MM, MX, MZ, NA, NG, NI, NO, NZ, OM, PO, PH, PL, PT, RO, RU, SC, ED, SS, SG, SK, SL, SM, SY, TJ, TM, TM, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
RNI BM, GH, GM, KE, LS, MM, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TM, AT, BE, BG, CH, CY, CZ, DE, DK, ER, RS, FT, FR, GB, GR, HU, IR, IS, IT, LT, LU, MC, NI, PL, PT, RO, SE, ST, SK, RB, PT, TR, BP, BJ, CF, CG, CI, CM, GA, GN, GQ, GM, ML, MR, NE, SN, TD, TO 20060406

PAGE 2-A

444579-17-39, 5,6,11,12,17,18-Hiexara-trinaphthylene-2,8,15-tricarboxylic acid triethyl ester 872140-78-89,
5,6,11,12,17,18-Hiexara-trinaphthylene-2,8,13-tricarboxylic acid triethyl ester 872140-78-99,
5,6,11,12,17,18-Hexara-trinaphthylene-2,8,13-tricarboxylic acid triethyl ester 872140-78-99, 5,6,11,12,17,18-Hexara-trinaphthylene-2,8,13-tricarboxylic acid tridodecyl ester 872140-80-39, 5,6,11,12,17,18-Hexara-trinaphthylene-2,8,13-tricarboxylic acid tridodecyl ester 872140-82-99, 5,6,11,12,17,18-Hexara-trinaphthylene-2,8,13-tricarboxylic acid tries(2,2,3,3,4,4,4-heptafluoro-butyl) ester 872140-83-99, 5,6,11,12,17,18-Hexara-trinaphthylene-2,8,13-tricarboxylic acid tries(2,2,3,3,4,4,4-heptafluoro-butyl) ester 872140-85-89, 5,6,11,12,17,18-Hexara-trinaphthylene-2,8,14-tricarboxylic acid tries(2-mathyl-butyl) ester 872140-86-89, 5,6,11,12,17,18-Hexara-trinaphthylene-2,8,14-tricarboxylic acid tries(2-mathyl-butyl) ester 872140-86-99 873140-89-19, 5,6,11,12,17,13-Hexara-trinaphthylene-2,8,15-tricarboxylic acid tries(2-mathyl-butyl) ester 872140-88-09 873140-89-19, 5,6,11,12,17,13-Hexara-trinaphthylene-2,8,15-tricarboxylic acid tribensylester 873140-80-09, 5,6,11,12,17,13-Hexara-trinaphthylene-2,8 tribenzylester RL: IMF (Industrial manufacture); TEM (Technical or engineered material uss); PREP (Preparation); USSS (Uses)

PRIORITY APPLN. INFO.:

Briefly described, embodiments of this disclosure include charge-transport materials (e.g., 2, 3, 8, 9, 14, 15-hexakiedodecylsulfanyl-5, 6, 12, 12, 17, 18-hexakezatrinaphthylene), methods of forming charge-transport materials, and methods of using the charge-transport materials. The charge-transport materials are used in organic electronic devices such as organic light-emitting diodes, lesers, photovoltaic cells, photodatectors, active and passive electronic devices, and memories.

IT 872140-83-59, 5,6,11,12,17,18-Hexakeza-trinaphthylene-2,8,15-tricarboxylic acid tripentafluorophenylenthyl ester 872140-84-69, 5,6,11,12,17,18-Hexakez-trinaphthylene-2,8,14-tricarboxylic acid tripentafluorophenylenthyl ester Briefly (Technical or engineered material use); PREP (Preparation); USES (Uses) (production of charge-transport materials containing hexakeztrinaphthylene for

organic electronic devices)
872140-83-5 CAPUS
Diquinoxalino[2,3-s-12',3'-c]phenazine-2,8,15-tricarboxylic acid,
tris[[pentafluorophenyl]methyl] ester [9CI] (CA INDEX NAME)

872140-84-6 CAPLUS Diquinoxalino(3,3-s12',3'-c)phenazine-2,8,14-tricarboxylic acid, tris[(pentafluorophenyl)acthyl) ester (9Cl) (CA INDEX NAME)

(production of charge-transport materials containing hexaezatrinaphthylene

organic electronic devices)
444579-17-3 CAPLUS
Diquinoxalino[2,3-e:2',3'-c]phenazine-2,8,15-tricarboxylic acid, triethyl
seter (9C1) (CA INDEX NAME)

872140-78-8 CAPLUS
Diquinoxalino[2,3-a:2',3'-c]phenazine-2,8,14-tricarboxylic acid, triethyl setc [9C1] (CA INDEX NAME)

872140-79-9 CAPLUS Diquinoxalino(2,3-s:2',3'-c)phenazine-2,8,15-tricarboxylic acid, tridodecyl ester (9CI) (CA INDEX NAME)

872140-80-2 CAPLUS
Diquinoxalino[2,3-a:2',3'-c]phenazine-2,8,14-tricarboxylic acid, tridodecyl ester (SCI) (CA INDEX NAME)

872140-81-3 CAPLUS
Diquinoxalino[2,3-a:2',3'-c]phenazine-2,8,15-tricarboxylic acid,
tris(2,2,3,3,4,4,4-heptafluorobutyl) ester (9CI) (CA INDEX NAMS)

PAGE 1-B

-- CF2- CF3

- cF2- CF3

RN 872140-82-4 CAPLUS
CN Diquinoxalino[2,3-a:2',3'-c)phenazine-2,8,14-tricarboxylic acid, tris[2,2,3,3,4,4,4-heptafluorobutyl) ester (9CI) (CA INDEX NAMS)

PAGE 1-B

-cF2-cF3

872140-85-7 CAPLUS
Diquinoxalino[2,3-a:2',3'-c]phenazine-2,8,15-tricarboxylic acid,
tris(2-methylbutyl) ester (9CI) (CA INDEX NAME)

872140-86-8 CAPLUS
Diquinoxalino[2,3-e:2',3'-c]phenazine-2,8,14-tricarboxylic acid,
tris(2-methylbutyl) ester (9CI) (CA INDEX NAME)

873140-87-9 CAPLUS Diquinoxalino[2,3-a:2',3'-c]phenazine-2,8,15-tricarboxylic acid, tris[2-(1-naphthalenyl)ethyl] ester (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

PAGE 3-A

872140-88-0 CAPLUS
Diquinoxalino[2,3-a:2',3'-c]phenazine-2,8,14-tricarboxylic acid,
tris[2-(1-naphthalenyl)ethyl] ester (9CI) (CA INDEX NAME)

872140-90-4 CAPLUS Diquinoxalino[2,3-e:2',3'-c]phenezine-2,8,14-tricarboxylic acid, tris(phenylmethyl) ester (9CI) (CA INDEX NAME)

L13 ANSWER 8 OF 181 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

AUTHOR (S):

CAPLUS COPYRIGHT 2006 ACS on STN
2005:1247836 CAPLUS
144:12838
144:128338
144:128338
164:128338
164:128338
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1

CORPORATE SOURCE: SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

872140-89-1 CAPLUS Diquinoxalino[3,3-e:2',3'-c]phenazine-2,8,15-tricarboxylic acid, tris(phenylmethyl) ester (9CI) (CA INDEX NAME)

OTHER SOURCE(S):

The synthesis of compds. I [R1 = 7-methoxybenzofuran-2-yl, quinoxalin-6-yl, 3-[2-pyrimidyl]phenyl, 5-[2-furyl]-3-pyrazolyl, etc.; R2 = 2-MeOC6H4, 2-benzimidszolyl, 5-methoxy-2-benzimoxazolyl, etc.], structurally related to the high-affinity dopamine D3 receptor ligand N-[4-(4-(2,3-dichlorophenyl)piperazin-1-yl]butyl]-7-methoxy-2-benzofurancarboxamide [II], is reported. All compds. were specifically designed as potential PET redioligands for brain D3 receptors visualization, having lipophilicity within a range for high brain uptake and weak nonspecific binding (2 < Clopp < 3.5) and bearing a methoxy substituent for easy access to labeling with the positron emitter isotope ilc. I [R1 = 4-(4-morpholinyl)phenyl, 4-(1-imidazolyl)phenyl, 5-(2-furyl)-3-pyrazolyl; R2 = 5-methoxy-2-benzisoxazolyl] displayed good D3 receptor affinities (Ki values 38.0, 22.6, and 21.3 nM, resp.) and were selective over D2 receptor. Moreover, these compds. were able to permeate the Caco-2 cell monolayer, differently from compound II. Although the goal to identify potential PET radioligands with subnanomolar affinities for D3 receptor was not achieved, the proposed strategy could be a starting point [737642-53-9] R16-63-2P
RL: PAC (Pharmacological activity); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (preparation, lipophilicity and brain dopamine D3 receptor binding affinities of N-(arylpiperazinyl) butyl heteroarylcarboxamides as potential positron emission tomog. ligands)
8-7064-55-9 CAPLUS
6-Quinoxalinecarboxamide, N-[4-(4-(5-methoxy-1,2-benzisoxazol-3-yl)-1-piperazinyl]butyl}- (9CI) (CA INDEX NAME)

873662-69-2 CAPLUS 6-Quinoxalimecarboxamide, N-[4-[4-(2-methoxyphenyl)-1-piperaxinyl]butyl]-(9CI) (CA INDEX NAME)

873662-82-9P 873662-89-6P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (preparation, lipophilicity and brain dopamine D3 receptor binding affinities of N-(arylpiperaxinyl)butyl heteroarylcarboxamides as potential positron emission tomog. ligands)
873662-82-9 CAPUIS
6-Quinoxalinecarboxamide, N-[4-(4-(5-methoxy-1,2-benzisoxazol-3-yl)-1-piperaxinyl)butyl}-, ethanedioate (1:2) (9CI) (CA INDEX NAME)

CM 1

CRN 873662-58-9 CMP C25 H28 N6 O3

CRN 144-62-7 CMF C2 H2 O4

HO-C-C-ON

873662-89-6 CAPLUS
6-Quinoxalinecarboxamide, N-[4-[4-(2-methoxyphenyl)-1-piperazinyl]butyl]-,
dihydrochloride [9CI] (CA INDEX NAME)

REFERENCE COUNT:

41 THERE ARE 41 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 9 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:1224419 CAPLUS
DOCUMENT NUMBER: 143:454349
TITLE: Proparation of quinoxalin-2-one derivatives as herbicide safeners
INVENTOR(S): Schaper, Wolfgang; Willms, Lothar; Rosinger, Christopher; Hacker, Erwin; Rose, Eckhard; Schmutzler, nir

Dirk

PATENT ASSIGNEE (S): SOURCE:

Dirk
Bayer Cropscience GmbH, Germany
U.S. Pat. Appl. Publ., 97 pp.
CODEN: USXXCO
Patent

DOCUMENT TYPE:

AUTHOR(S):

Hexaezetrinaphthylene Derivative

Kaefareni, Bilal R.; Kondo, Takeshi; Yu, Junsheng;
Zhang, Ging; Dattilo, Davide: Risko, Chad; Jonee,
Simon C.; Barlow, Stephen; Domercq, Benoti; Amy,
Fabrice; Kahn, Antoine: Bredas, Jean-Luc; Kippelen,
Bernard; Marder, Seth R.

CORPORATE SOURCE:

Center for Organic Photonics and Electronics (COPE),
School of Chemistry and Biochemistry and School of
Electrical and Computer Engineering, Georgia Institute
of Technology, Atlanta, GA, 3012, USA

SOURCE:

Journal of the American Chemical Society (2005),
127(47), 16158-16159

CODEN: JACSAT; ISSN: 0002-7863

American Chemical Society

DOCUMENT TYPE:
Journal
LANGUAGS:
Biglish

AB An isomeric mixture of a tris(pentafluorobenzyl ester) derivative of
hexaszatrinaphthylene forms etable amorphous films with an effective
charge-carrier mobility of 0.02 cm2/Vs, while the pure 2,8,15-isomer
exhibits videly differing morphologies and carrier mobilities (0.001-0.07

cm2/Vs), depending critically on the processing conditions.

T 87214-0-8-15 972140-84-59

RL; DEV (Device component use); PRP (Properties); SPN (Synthetic
preparation); PRSP (Preparation); USSS (Uses)
(carrier mobility in amorphous hexaszatrinaphthylene derivative)

RN 87214-0-8-15 CAPUS

Diquinoxalino[2,3-a:2',3'-c]phenszine-2,6,15-tricerboxylic acid,
tris[(pentafluorophenyl)methyl] ester (9CI) (CA INDEX NAME)

872140-84-6 CAPLUS Diquinoxalino[2,3-e:2',3'-c]phenazine-2,8,14-tricarboxylic acid, triel[pentalluorophenyl]methyl] ester (9CI) (CA INDEX NAME)

PAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

| PAT | TENT | NO. | | | KIN | D | DATE | | | APPL | ICAT | ION : | NO. | | D | ATE | | |
|-----|------|------|------|-----|-----|-----|------|------|-----|------|------|-------|------|------|-----|------|-----|----|
| | | | | | | - | | | | | | | | | - | | | |
| us | 2005 | 2560 | 00 | | A1 | | 2005 | 1117 | 1 | US 2 | 005- | 1270 | 16 | | 2 | 0050 | 511 | |
| DE | 1020 | 0402 | 3332 | | A1 | | 2006 | 0119 | | DE 2 | 004- | 1020 | 0402 | 3332 | 2 | 0040 | 512 | |
| MO | 2005 | 1126 | 30 | | A1 | | 2005 | 1201 | | NO 2 | 005- | EP44 | 45 | | 2 | 0050 | 426 | |
| | w: | AB. | AG. | AL. | AM. | AT. | AU. | AZ. | BA. | BB. | BG. | BR. | BW, | BY. | BZ. | CA. | CH, | |
| | | CN. | co, | CR, | CU, | CZ, | DK, | DM. | DZ, | EC. | EE, | EG, | ES, | FI. | GB, | GD, | GE, | |
| | | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG, | KM, | KP, | KR, | KZ, | LC, | |
| | | LK, | LR, | LS. | LT. | LU, | LV. | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NA, | NI, | |
| | | NO, | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SΚ, | SL, | SM, | |
| | | SY, | TJ. | TM, | TN, | TR, | TT. | TZ. | UA. | υa, | US, | υz, | VC, | VN, | Yυ, | ZA, | ZM, | ZW |
| | RW: | BW, | GH, | GM, | KE, | LS, | MW, | MZ, | NA, | SD. | SL, | SZ. | TZ, | UG, | ZM, | ZW, | AM, | |
| | | AZ, | BY, | KG, | KZ, | MD, | RU, | TJ, | TM, | AT, | BE, | BO, | CH, | CY, | CZ, | DE. | DK, | |
| | | EE, | ES, | FI. | PR. | GB, | GR, | HU, | IE, | IS, | IT, | LT, | LU, | MC, | NL, | PL, | PT, | |
| | | RO, | SE, | SI, | SK, | TR, | BF, | BJ, | CF. | CG, | CI, | CM, | GA, | GN, | GQ, | GW, | ML, | |
| | | | | | | | | | | | | | | | | | | |

MU. SE, SI, SK, TR, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO:
OTHER SOURCE(S): MARPAT
GI DE 2004-102004023332A 20040512

$$Y_n = \bigvee_{N=0}^{R^1} X$$

The quinoxalin-2-one derivs. I [X = O or S; Y = halo, cyano, nitro, alkyl, alkenyl, alkynyl, etc.; n = 0, 1, 2, 3 or 4; Rl = H, OH, NH2, alkylamino, dialkylamino, (un)substituted alkyl, alkenyl, alkynyl or alkoxy, cycloalkyl, cycloalkenyl, aryl or heterocyclyl; R2 = H or (un)substituted alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl or heterocyclyl; or I selts are prepared as herbicide safeners.
865312-47-00
RH. AGR (Agricultural use); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation as herbicide safener)
869312-47-0 CAPLUS
6-Quinoxalinecarboxylic acid, 1,2-dihydro-2-oxo-3-(2-thienyl)-, ethyl ester (SCI) (CA INDEX NAME)

L13 ANSWER 10 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:1184873 CAPLUS
DOCUMENT NUMBER: 144:98799
TITLE: High Charge-Carrier Mobility in an Amorphous

444579-17-3 872140-78-8
RL: PRP (Properties)
(carrier mobility in amorphous hexaszatrinaphthylene derivative)
444579-17-3 CAPUS
Diquinoxalino(2,3-e.2',3'-c)phenazine-2,8,15-tricerboxylic acid, triethyleater (9CI) (CA INDEX NAME)

872140-78-8 CAPLUS Diquinoxalino[2,3-a:2',3'-c]phenazine-2,8,14-tricarboxylic acid, triethyl ester (9CI) (CA INDEX NAME)

THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 11 OF 181 ACCESSION NUMBER:

CAPLUS COPYRIGHT 2006 ACS on STN 2005:1050874 CAPLUS

DOCUMENT NUMBER: TITLE:

2005:1050874 CAPLUS
143:326207
Preparation and pharmaceutical compositions of
pyrrolidine derivatives as inhibitors of dipeptidyl
peptidase-iv (DPP-iv)
Akritopoulou-Zanze, Irin; Darczak, Daria; Dinges,
Jurgen; Djuric, Stevan W.; Hoff, Ethan D.; Kopecka,
Hana A.; Patel, Jyoti R.; Pei, Zhonghua; Shuai, Oi;
Sarris, Kathy; Sham, Hing L.; Wiedeman, Paul E.
USA

PATENT ASSIGNEE(S): SOURCE:

U.S. Pat. Appl. Publ., 51 pp. CODEN: USXXCO Patent

DOCUMENT TYPE

English 1 PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. US 2005215603
PRIORITY APPLM. INFO.:
OTHER SOURCE(S):
GI

KIND DATE A1 20050929

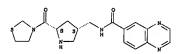
MARPAT 143:326207

APPLICATION NO. US 2004-795622 US 2004-795622

20040308 20040308

Title compds. I [R1 = aryl, alkyl, cycloalkyl, etc.; D = CO, O, SO2, CONH, etc.; L = bond, -CH2-, aryl, etc.; A = CO, NHSO2, NHCO, etc.; X = CHF, CH3, O, S, etc.], and their pharmaceutically acceptable salts, are prepared and disclosed as inhibitors of dispeptidyl peptidese ethics. The control of the cont

(Uses) (drug candidate; preparation of pyrrolidine derivs. as inhibitors of dispetidyl peptidase-iv (DDP-iv) (DS-iv) (



5296-56-6 CAPLUS Quinoxalinecarboxamide, N-[{(38,58)-5-(1-pyrrolidinylcarbonyl)-3-rrolidinyl|methyl1-(9CI) (CA INDEX NAME)

L13 ANSWER 12 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:1028081 CAPLUS
DOCUMENT NUMBER: 143:132602
TITLE: Preparation and pharmaceutical compositions of pyrrolidine derivatives as inhibitors of dipeptidyl peptidase-iv (OPP-iv)
Akritopoulou-Zanus, Irini; Darczak, Daria; Dinges, Jurgen; Djuric, Stevan W.; Hoff, Ethan D.; Kopecke, Hans A.; Patel, Jyoti R.; Pei Zhonghus, Shusi, Qi; Sarris, Kathy; Shas, Hing L.; Wiedeman, Paul E.

USA

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. PATENT NO. KIND DATE US 2005209249
PRIORITY APPLN. INE A1 20050922 US 2005-75319 US 2004-551079P INFO.: MARPAT 143:326202

20050308 P 20040308

U.S. Pat. Appl. Publ., 50 pp. CODEN: USXXCO

Patent English

Absolute stereochemistry.

(Uses) (drug candidate; preparation of pyrrolidine derivs. as inhibitors of dispeptidyl peptidase-iv (DDP-iv)) 853294-89-8 CAPLUS 6-Quinoxalinecarboxamide, N-[[(38,58)-5-(3-thiszolidinylcarbonyl)-3-pyrrolidinyllanthyll- (9CI) (CA INDEX NAME)

865296-56-6 CAPLUS 6-Quinoxalinecarboxamide, N-[[(35,58)-5-(1-pyrrolidinylcarbonyl)-3-pyrrolidinyl]methyll- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

ACCESSION NUMBER:

ACCESSION NUMBER:

DOCUMENT NUMBER:

AS APPROACH TIS COPPRIGHT 2006 ACS on STN

2005:989948 CAPLUS

143:433305

APPROACH TIS COMPANY OF THE APPROACH TO THE

complex-tethered short fluorescent DNA probes to human telomere repetitive DNA)

ROM 863942-78-7 CAPUUS

NA Adenosine, 2'-deoxy-5'-0-[[[6-[(dipyrido[3,2-a:2',3'-c]phenazin-11-ylcarbony]) mainto | hexyll oxyl hydroxyphosphinyll cytidylyl-(3' →5')-2'-deoxycytidylyl-(3', -5')-2'-thynidylyl-(3', -5')-1'-thynidylyl-(3'-5')-1'-thynidy

Absolute stereochemistry.

PAGE 1-B

PAGE 2-A

868744-57-4 868744-70-1 868745-17-9

RL: ARG (Analytical reagent use): BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(asym. cooperativity in tandem hybridization of enantiomeric metal complex-techred short fluorescent DNA probes)

868744-57-4 CAPULS

Ruthenate(4-), [5'-0-[[[6-{[(dipyrido[3,2-s-2',3'-c]phenazin-11-yl-KN4,KN5)carbonyl]maino]hexyl]oxyl)hydroxyphosphinyl]-2'-deoxyyetylyl;1-(3'-5')-2'-deoxyyetyldylyl-(3'-5')-2'-deoxyyetyldylyl-(3'-5')-2'-deoxyyethylyl-(3'-5')-2'-deoxyyethylyl-(3'-5')-2'-deoxyyethylyl-(3'-5')-2'-deoxyyethylyl-(3'-5')-2'-deoxyyethylyl-(3'-5')-2'-deoxyyethylyl-(3'-5')-2'-deoxyyethylyl-(3'-5')-2'-deoxyyethylyl-(3'-5')-2'-deoxyyethylyl-(3'-5')-2'-deoxyyethyl-(3'-5')-2'-deoxyyethylyl-(3'-5')-2'-deoxyyethyl-(3'-5')-2'-deoxyethyl-(3'-5')-2'-deoxyyethyl-(3'-5')-2'-deoxyethyl-(3'-5')-2'-deo

PAGE 1-B

PAGE 2-A

●5 H*

RN 868744-70-1 CAPLUS

RN Ruthenate(4-), [5'-0-{[[6-[[(dipyrido[3,2-a:2',3'-c]phenazin-11-y1-xN4,xN5]carbony]lasino[hexyl]cxy]hydroxyphosphinyl]-2'-deoxycytidylyl-[3'-s')-2'-deoxycytidylyl-[3'-s')-2'-deoxycytidylyl-[3'-s')-2'-deoxycytidylyl-[3'-s')-2'-deoxydenylyl-[3'-s')-2'-deoxydenylyl-[3'-s')-2'-deoxydenylyl-[3'-s')-2'-deoxydenylyl-[3'-s']-2'-deoxydenylyl-[3'-s

PAGE 1-A

PAGE 2-B

868745-17-9 CAPLUS
Ruthenate(4-), [5'-o-{[[6-[[(dipyrido[3, 2-e:2', 3'-c]phenazin-11-yl-tN4, KN5]carbonyl]asino]hexyl]oxyl hydroxyphosphinyl]-2'-deoxycytidylyl-(3'-s')-2'-deoxycytidylyl-(3'-s')-2'-deoxycytidylyl-(3'-s')-2'-deoxydenylyl-(3'-s')-2'-deoxydenylyl-(3'-s')-2'-deoxydenylyl-(3'-s')-2'-deoxydenylyl-(3'-s')-2'-deoxydenylyl-(3'-s')-2'-deoxydenylyl-(3'-s')-2'-deoxydenylyl-(3'-s')-2'-deoxydenylyl-(3'-s')-2'-deoxydenylyl-(3'-s')-2'-deoxydenosinato(6-)]bie(1,10-phenanthroline-KN1,KN10-), pentahydrogen, (OC-6-33-A)- (9CI) (CA

PAGE 1-A

PAGE 2-A

●5 H+

PAGE 2-A

●5 H*

THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

PERFERENCE COUNT.

L13 ANSWER 14 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:800492 CAPLUS
105:800492 CAPLUS
141:386895
A phase-switch purification approach for the expedient removal of tagged reegents and scavengers following their application in organic synthesis Siu, Jeson; Baxendale, Ian R.; Lewthwaite, Russell A.; Ley, Steven V.
CORPORATE SOURCE: Depertment of Chemistry, University of Cambridge, Cambridge, CSS 1EW, UK

Department of Chemistry, University of Cambridge Cambridge, C93 1EM, UK Orgenic & Biomolecular Chemistry (2005), 3(17), 3140-3160 CODEN: OBCRAK; ISSN: 1477-0520 Rayel Society of Chemistry Journal

UBLISHER: OCUMENT TYPE:

DUBLISHER: Royal Society of Chemiatry

DOUMENT TYPE: Bournal

LANGULAGE: Regish

AB In this paper a variety of expedient chemical transformations and

purifications achieved via a generic catch and release methodol., based on
a synthetically inert bipyridyl chelating tag that can be selectively
captured with a resin-bound copper(II) species, were reported. Utilizing
this approach it was possible to derive many of the same benefits associated
with both aclid phase synthesis and supported reagent methods.

IT 866789-73-59

RI: PUR (Purification or recovery); SPN (Synthetic preparation); PREP

RI: PUR (Purification or recovery); SPN (Synthetic preparation); PREP

(Preparation) (preparation) desides using amines and carboxylic acid es reactants and
N-(cyclohexylcarbonimidoyl)bipyridine amine as coupling egent and study
of phase-switch purification approach for expedient removal of tagged
resgents and seavengers)

RN 865789-75-5 CAPLUS

CN 6-Quinoxalinecerboxamide, N-[(1,3,5-trimethyl-1H-pyrazol-4-yl)methyl](9CI) (CA INDEX NAME)

REFERENCE COUNT:

61 THERE ARE 61 CITED REFERENCES AVAILABLE FOR THIS

853914-90-6 CAPLUS
Copper(1+), (ethyl dipyrido(3,2-e:2',3'-c)phenezine-ll-carboxyleteKM4,KM5)bie(triphenylphosphine)-, (T-4)-, tetrefluoroborate(1) (9C1) (CA INDEX NAME)

853914-89-3 C57 H44 Cu N4 O2 P2 CC8

CRN 14874-70-5 CMF B F4 CCI CCS

862288-29-7
RL: PRP (Properties)
(Raman spectroscopy and DPT calcns. in study of ground- and excited states of Cu(1) and Re(1) complexes with dipyridophenasine ligands)
852288-29-7 CAPLUS
Rhenate(1-), tricarbonylchloro(ethyl dipyrido(3,2-a:2',3'-c)phenasine-ll-carboxylete-kMe, KMS)-, (OC-6-44)- (9CI) (CA INDEX NAME)

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ACCESSION NUMBER:
DOCUMENT NUMBER:
105:504199 CAPLUS
143:219327
Photoexcitation in Cu(I) and Re(I) Complexes
Containing Substituted Dipyrido(3, 2-e; 2', 3'c]phenszine: A Spectroscopic and Density Punctional
Theoretical Study
Malsh, Penny J.; Gordon, Keith C.; Lundin, Natasha J.;
Blackman, Allan G.
CORPORATE SOURCE:
50URCE:
50URC

LISHER:

American Chemical Bociety

JOHN 1 TYPE:

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JOHN 2 John 1 Lisher:

LISHER:

American Chemical Bociety

JOHN 2 John 1 Lisher:

LISHER:

JOHN 2 J LANGUAGE:

853914-87-1 RL: PRP (Properties)

(ligend; Raman spectroacopy and DFT calens, in study of ground-accited states of Cu(I) and Re(I) complexes with dipyridophenazine

ligands)
853914-87-1 CAPLUS
Dipyrido[3,2-a:2',3'-c]phenazine-ll-carboxylic acid, ethyl ester (9CI)
(CA INDEX NAME)

REFERENCE COUNT:

CORPORATE SOURCE:

THERE ARE 49 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 16 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2005:339438 CAPLUS DOCUMENT NUMBER: 143:70403 Complexes of Street, 100 Com

143:70403
Complexes of Functionalized Dipyrido[3,2-a:2',3'-c]phenazine: A Synthetic, Spectroscopic, Structural, and
Density Functioned Theory Study
Lundin, Natasha J.; Malsh, Penny J.; Howell, Sarah L.;
McGarvey, John J.; Blackman, Allan G.; Gordon, Keith
C. AUTHOR (S)

C.
Department of Chemistry, MacDiarmid Institute for Advanced Materials and Nenotechnology, University of Otago, Dunedin, N. Z.
Inorganic Chemistry (2005), 44(10), 3551-3560
CODEN: INOCAJ: ISSN: 0020-1669
American Chemical Society
Journal

PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): English CASREACT 143:70403

R SOURCE(S): CASERACT 143:70403
The ligands 11-bromodipyrido[3,2-a:2',3'-c]phenazine and St
dipyrido[3,2-a:2',3'-c]phenazine-11-carboxylate were prepared and
coordinated to Ru(II), Re(I), and Cu(I) metal centers. The electronic
effects of substitution of dipyrido[2,3-a:3',2'-c]phenazine (dpps) were
atudied by spectroacopy and electrochem, and some photophys. properties
were studied. The crystal structures of [Re(I)(CO)3CI] (I = Et
dipyrido[3,2-a:2',3'-c]phenazine-11-cerboxylate or 11-bromodipyrido[3,2a:2',3'-c]phenazine) are presented. D. functional theory calcus. on the
complexes show only small deviations in bond lengths and angles (most
bonds within 0.02 Å, most angles within 2*) from the

crystallog, data. Also, the vibrational spectra of the strongest Raman and IR bands are predicted to within an average 6 cm-1 for [Re(L)|(CO)3C1] and [Cu(L)|(triphenylphosphine)2]BF4 (in the 1000-1700 cm-1 region).

Spectroscopic and electrochem. evidence suggest that reduction of the complex causes structural changes across the entire dppz ligand. This is unusual as dppz-based ligands typically have electrochem, properties that suggest charge localization with reduction on the phenazine portion of the ligand. The excited-state lifetimes of the complexes were measured, and they range from ca. 200 Ns for the [Ru(L)[2,2"-bhpyridine)2](PF6)2 complexes to over 2 µs for [Cu(11-bromodipyrido], 2-a-2', 3'-c')phenazine|(PP6)2] [SP8] at room temperature The emission spectra suggest that the unusually long-lived excited states of the Cu complexes result from metal-to-ligand charge transfer (MLCT) transitions as they are completely quenched in MeOH. Slectroluminescent (ilms may be fabricated from these compde; they show MLCT state emission even at low doping levels [co.1% by weight in Phylical-space] process); PRP (Properties); SPN (Synthatic preparation); TEM (Technical or engineered material usel); PRED (Properation); PRDC (Process); USES (Uses) (preparation, crystal structure, exptl. and calculated mol. structure, fluorescence, reduction potentials, vibrational spectra and application in electroluminescent film) 757350-96-9 CAPLUS

Rhenium, tricarbonylchloro(ethyl dipyrido[3,2-a:2',3'-c]phenazine-11-carboxylate-kN4,kN5)-, (OC-6-44)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

853914-87-1P
RL: CPS (Chemical process); PEP (Physical, engineering or chemical process); RPP (Properties); RCT (Reactant); SPN (Synthetic preparation); PRDP (Properation); PRDP (Process); RACT (Reactant or reagent) (preparation, mol. structure from DPT calcus., fluorescence, reduction potentials and complexation with copper, rhenium and ruthenium) 853914-87-1 CAPUIS Dipyrido(3,2-s:2',3'-c]phenazine-11-carboxylic acid, ethyl ester (9CI) (CA INDEX NAME) IT

853914-91-7P 853914-95-1P RL: CPS (Chemical process); PEP (Physical, engineering or chemical

2

16919-18-9 F6 P CCS

REFERENCE COUNT:

THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 17 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:324138 CAPLUS
DOCUMENT NUMBER: 142:392428
ITITLE: Proparation of heterocyclic compounds as antifungal agents
INVENTOR(6): Nakamoto, Kazutaka; Tsukada, Itaru; Tanaka, Keigo; Matsukura, Masayuki; Haneda, Toru; Inoue, Satoshi; Ueda, Norihiro; Abe, Shinya; Hata, Katsura; Watanabe, Haoaki
PATENT ASSIGNEE(8): Siesi Co., Ltd., Japan
PCT Int. Appl., 418 pp.
CODEN: PIXENZ
LANGUAGE: Patent
LANGUAGE: Japanese

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.

KIND DATE

APPLICATION NO.

DATE

process); PRP (Properties); SPN (Synthetic preparation); PREP (Preparation); PROC (Process) (preparation), PROC (Process) (preparation, mol. structure from DPT calons., fluorescence, reduction potentials of) 853914-91-7 CAPLUS (Synthetics) CRN 853914-90-6 CMP C57 H44 Cu N4 O2 P2 . B F4 CM 2 CRN 853914-89-3 CMF C57 H44 Cu N4 O2 P2 CCI CCS

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JP 2003-342273
JP 2004-68186
JP 2004-232617
                                                     20030930
20040310
20040809
PRIORITY APPLA
                                                   AAAAA
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OTHER SOURCE(S): MARPAT 142:392428

The title compds., e.g. I [ring Al is optionally substituted 3-pyridyl, optionally substituted quinolyl, etc.; Xl is NNCO, etc., and ring E is furyl, thienyl, pyrrolyl, Ph, pyridyl, tetrazolyl, thiszolyl, or pyrazolyl; provided that Al may have one to three substituents and E has one or two substituents], are prepared 2.6-Diamino-N-(5-(4-fluorophenoxy)furan-2-ylmethyl)nicotinamide was prepared in a multistep process. Compds. of this invention in vitro showed MIC values of 0.1 µg/mL to 6.25 µg/mL against Candida.
849810-87-3P
RE: PAC (Pharmacological activity); SPN (Synthetic preparation); TMU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of heterocyclic compds. as antifungal agents) 849810-87-3 CADUE 6-Ouinoxalinecarboxamide, N-[(3-phenoxyphenyl)methyl]-, momo(trifluoroacotate) (9CI) (CA INDEX NAME)

CH 1

CRN 849810-86-2 CMF C22 H17 N3 O2

CRN 76-05-1 CMF C2 H F3 O2

ARE 56 CITED REPERENCES AVAILABLE FOR THE RECORD. ALL CITATIONS AVAILABLE IN THE RE PORUG.

1 CAPLUS COPYRIGHT 2006 ACS on STN
2005:324132 CAPLUS
142:32427
Preparation of N-heterocyclyl amides and sulfonamides as p38 kinase inhibitors
Dugar, Sundeep; McEnroe, Glen
Sclos Inc., USA
PCT Int. Appl., 195 pp.
CUDEN: PIXXD2
Patent
Bilsh
1 REFERENCE COUNT:

L13 ANSWER 18 OF 181 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| TENT | INFOR | ITAM | ON: | | | | | | | | | | | | | | |
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| w | 0 2005 | 0330 | 72 | | A2 | | 2005 | 0414 | | WO 2 | 004- | US32 | 403 | | 2 | 0040 | 930 |
| w | 0 2005 | 0330 | 72 | | A3 | | 2006 | 0112 | | | | | | | | | |
| | W: | AE, | AG. | AL. | AM. | AT. | AU. | AZ. | BA. | BB. | BG. | BR. | BW. | BY. | BZ. | CA. | CH. |
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| TORT | TY APE | | | | | | | | | 118 2 | 003- | 5076 | 330 | | P 2 | 0030 | 930 |
| | SOURCE | | | | MAR | PAT | 142: | 3924 | | | | | | | _ | | |

849748-71-6 CAPLUS 6-Quinoxalinecarboxamide, N-[2-[[(1S)-1-phenylethyl]amino]-4-pyrimidinyl]-N-(4-piperidinylmethyl)- (9CI) (CA INDEX NAME)

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE US 2005032805 US 6849660 20050210 20050201 US 2004-914256 US 2000-630122 20040809

The title compde. I [R1 = alkyl, cycloslkyl, heterocycloslkyl, aryl; L = CO, EO2; X = O, CO, (un) substituted CH2, NH; n = 0-1; R2 = H, alkyl, aryl, etc.; Y = (un) substituted NH2, OR; one of 21 and 22 = CH, and the other is either CH or N], useful for inhibiting pl8 kinses, were prepared E.g., a multi-step synthesis of (18)-II, etarting from 4-amino-2-chloropyridine and 2-naphthoyl chloride, was given. The compde. I were tested against pl8c kinses in the diluted whole blood assay (bold data were given for representative compds. I). The pharmacoutical composition comprising the compound I is disclosed.
849746-04-9P 849748-62-5P 849746-71-6P
RL; PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapoutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(Uses)
(preparation of N-heterocyclyl amides and sulfonamides as pl8 kinase inhibitors)
48746-04-9 CAPLUS
6-Ouinoxalinecarboxamide, N-ethyl-N-[2-{{13}-1-phenylethyl}amino]-4-pyrimidinyl]- (9CI) (CA INDEX RAME)

Absolute stereochemistry.

849748-62-5 CAPLUS
1-Piperidinecarboxylic acid, 4-[[[2-[[(1S)-1-phenylethyl]amino]-4-pyrimidinyl[6-quinoxalinylcarbonyl]amino]methyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

PRIORITY APPLN. INFO:

OTHER SOURCE(S):

AB Provided are antibacterial biaryl compds. having micromolar MIC activity against Gran-neg and Oran-pos. pathogens, including a methicillin-resistant B. aureus strain. Other embodiments of invention are methods of treating bacterial infection in a mammal by administering to the mammal an effective amount of a compound described herein. The inhibitory effect of some of the compds. on bacterial translation was determined

IT 797770-86-6

RI: PAC (Pharmacological activity); TRU (Therapeutic use); BIOL (Biological study); USES (Uses)

(antibacterial biaryl compds. in relation to inhibiting bacterial translation and overcoming methicillin resistance)

RN 797770-86-6 CAPLUS

CN 6-Quinoxalinecerboxamide, N-{1-{4-{3-(dimethylamino)propoxylphenyl}-2-oxoneric complex companies of the companies o 797770-88-6
6-Quinoxalinecarboxamide, N-[1-[4-[3-(dimethylamino)propoxy]phenyl]-2-oxo2-(1-piperazinyl)ethyl]-1,2,3,4-tetrahydro-2,3-dioxo- (9CI) (CA INDEX

Me2N- (CH2) 3

L13 ANSWER 20 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER: 1005;99161 CAPLUS
DOCUMENT NUMBER: 142:13802 Preparation of quinoxalinecarboxamides as antivirals
ARBOYUN: Rong, Frank; Mu, Jim; Harris, Clayton;
Chow, Suetying
USA
USA
USA
USA
USA
USA
PARENT TYPE: Patent
EANGUAGE: Patent
EANGUAGE: PATENT INFORMATION:
1 APPLICANTS

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APPLICANTS APPLICANT DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE

APPEICATION NO. US 2004-826439 US 2003-463257P US 2005026923
PRIORITY APPLN. INFO.:
OTHER SOURCE(S):
GI A1 20050203 6/8261 MARPAT 142:198008

Title compds. [I; Z = NH, O; X = OH, NH2, OR, NHR, NR2, SR; R1, R2 = H, (substituted) alkyl, alkenyl, alkynyl, aryl, fused aryl, heterocyclyl, fused heterocyclyl; R1R2 = atoms to form a 5-6 membered ring; R3 = (substituted) alkyl, alkenyl, alkynyl, aryl, fused aryl, heterocyclyl, fused heterocyclyl; R, R4 = H, (substituted) alkyl, alkenyl, alkynyl, aryl, fused heterocyclyl; Rused heterocyclyl, vere prepared Thus, title compound (II), (preparation from L-5-hydroxytryptophan, -diminobenzoic acid, and 4-fluorobenzyl given) showed inhibitory activity with IC50 <10 µM in an hepatitis C virus (RCV) NSSB replicon assay.

835922-72-0P 835922-73-1P 835922-74-2P 835922-76-8P 835922-76-8P 835922-79-9 RJS 835922-80-8P 835922-80-80 PS 835922-80-80 PS 835922-80-80 PS 83592-80-80 PS 835922-80-80 PS 835922-80 PS 835922-

Absolute stereochemistry.

835922-73-1 CAPLUS L-Tryptophan, N-[12,3-bis(4-fluorophenyl)-6-quinoxalinyl]cerbonyl]-5-hydroxy-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

835922-79-7 CAPLUS 6-Quinoxalinecarboxamide, N-[(18)-2-amino-1-(1H-indol-3-ylmethyl)-2-oxoethyl)-2,3-bis(4-fluorophenyl)- (9CI) (CA INDEX NAME)

835922-81-1 CAPLUS L-Tryptophan, N-(2-phenazinylcarbonyl)-, methyl ester (9CI) (CA INDEX NAMS)

Absolute stereochemistry.

835922-82-2 CAPLUS L-Tryptophan, N-(2-phenazinylcarbonyl)- (9CI) (CA INDEX NAME) Absolute stereochemistry.

835922-74-2 CAPLUS L-Tryptophan, N-[[2,3-bis(4-fluorophenyl)-6-quinoxalinyl]csrbonyl]-5-hydroxy- [9C1) (CA INDEX NAME)

835922-76-4 CAPLUS L-Tryptophan, N-[[2,3-bis(4-fluorophenyl)-6-quinoxalinyl]carbonyl]- (9CI) (CA INDEX IAME)

835932-78-6 CAPLUS L-Tryptophan, N-{2,3-bis{4-fluorophenyl}-6-quinoxalinyl}carbonyl}-, methyl ester (9C1) (CA INDEX NAME)

MPCT CAMES

835922-95-7 CAPLUS L-Tryptophan, N-[(2-cyclohexyl-3-phenyl-6-quinoxalinyl)carbonyl]-5-hydroxy-, methyl eater (SCI) (CA INDEX NAME)

835922-96-8 CAPLUS L-Tryptophan, N-{(2-cyclohexyl-3-phenyl-6-quinoxalinyl)carbonyl}-5-hydroxy-(9CI) (CA INDEX NAME)

835922-97-9 CAPLUS L-Tryptophan, N-[(3-cyclohexyl-2-phanyl-6-quinoxalinyl)carbonyl)-5-hydroxy-nethyl ester (9C1) (CA INDEX NAMS)

Absolute stereochemistry.

835922-98-0 CAPLUS L-Tryptophan, N-[(3-cyclohexyl-2-phenyl-6-quinoxalinyl)carbonyl]-5-hydroxy-(9C1) (CA INDEX NAME)

835922-83-3 835922-84-4 835922-85-5
835922-86-6 835922-87-7 835922-89-9
835922-90-2 835922-92-4
RL: PAC (Phermacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (preparation of quinoxalinecarboxamides as antivirals)
835922-83-3 CAPLUS
L-Tryptophan, N-[(2,3-diphenyl-6-quinoxalinyl)carbonyl)- (9CI) (CA INDEX NAME)

835922-84-4 CAPLUS L-Tryptophan, N- (2,3-diphenyl-6-quinoxalinyl)carbonyl]-5-hydroxy- (9CI) (CA INDEX RAME)

835922-85-5 CAPLUS L-Tryptophan, N-[(2.3-diphenyl-6-quinoxalinyl)carbonyl]-5-hydroxy-, methyl ester (9C1) (CA INDEX NAME)

Absolute stereochemistry.

835922-86-6 CAPLUS L-Tryptophan, N-{2,3-di-2-pyridinyl-6-quinoxalinyl}carbonyl}-, methyl eater (9C1) (CA INDEX NAME)

solute stereochemistry.

835922-87-7 CAPLUS L-Tryptophan, N-{(2,3-di-2-pyridinyl-6-quinoxalinyl)carbonyl]- (9CI) (CA INDEX NAWL)

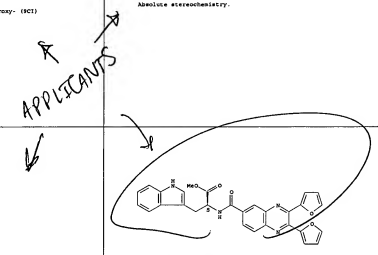
Absolute stereochemistry.

835922-89-9 CAPLUS L-Tryptophan, N-{{2-(4-chlorophenyl)-3-(4-methylphenyl)-6-quinoxalinyl]earbonyl}-5-hydroxy- (9CI) (CA INDEX NAME)

815922-90-2 CAPLUS L-Tryptophan, N-[{2-(4-chlorophenyl)-3-(4-methylphenyl)-6-quinoxalinyl]carbonyl]-5-hydroxy-, methyl ester (9CI) (CA INDEX NAME)

835922-92-4 CAPLUS L-Tryptophan, N-([2,3-di-2-furany1-6-quinoxaliny1)carbony1]-, methyl ester (9C1) [CA INDEX NAME]

Absolute stereochemistry.



L13 ANSWER 21 OF 181
ACCESSION NUMBER:
DOCUMENT NUMBER:
171TLE:
1NVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NORTH.
FAMILY ACC. NORTH.
PATENT LORDING. COURT:
FAMILY ACC. NORTH.
PATENT LORDING. COURT:
FAMILY ACC. NORTH.
FAMILY A

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PA | TENT | NO. | | | KIN | D | DATE | | | APPL | ICAT | ION | NO. | | D. | ATE | |
|----|------|------|-----|-----|-----|-----|------|------|-----|------|-------|------|-----|-----|-----|------|-----|
| | | | | | | - | | | | | | | | | - | | |
| | 2005 | | | | A2 | | | 0127 | | WO 2 | 004 - | US21 | 634 | | 2 | 0040 | 709 |
| WO | 2005 | 0070 | 99 | | A3 | | 2005 | 0414 | | | | | | | | | |
| | ₩: | AE, | AG, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BR, | BW, | BY, | BZ, | CA, | CH, |
| | | CN, | œ, | CR, | CU, | CZ, | DE. | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FI, | GB, | GD, |
| | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG, | KP, | KR, | KZ, | LC, |
| | | LK, | LR. | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NA, | NI, |
| | | NO. | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | SY, |
| | | TJ, | TM, | TN, | TR, | TT, | TZ, | UA, | UG, | US, | υZ, | VC. | VN, | Yυ, | ZA, | ZM, | ZW |
| | RW: | BW, | GH, | GM, | KE, | LS, | MW, | MZ, | NA, | SD, | SL, | SZ, | TZ, | υσ, | ZM, | ZW, | AM, |
| | | AZ, | BY, | KG, | ĸz, | MD, | RU, | TJ, | TM, | AT, | BE. | BG, | CH, | CY, | CZ, | DE, | DK, |
| | | EE, | ES, | PI, | PR, | GB, | GR, | HU, | IE, | IT, | LU, | MC, | NL, | PL, | PT, | RO, | SE, |
| | | SI, | SK, | TR, | BF, | BJ, | CF, | CG, | ÇI, | CM, | GA, | GN, | GQ, | GW, | ML, | MR, | NE, |
| | | | | | | | | | | | | | | | | | |

SI. SK. TR. SN. TD. TG PRIORITY APPLN. INFO.: OTHER SOURCE(S): GI US 2003-486339P MARPAT 142:176856 P 20030710

Title compds, represented by the formula I [wherein X - N or C; Ri, R2 = independently H, (cyclo]alkyl, alkoxy, heterocycly(alkyl), (heterolaryl, (heterolaryl, can) substituted amino; R3-R6 = independently H, cyano, (heterolaryl, (cyclo]alkyl, etc., with a provise) were prepared as PKB inhibitors. Por example, reaction of 4,5-diaminopyrimidine with

2.2'-thenyl gave II in 19% yield. I were tested for inhibition of PKB in PKBa, PKBB and PKBy in vitro kinese assay. Thus, I and their pharmaceutical compns. are useful as PKB inhibitors for the treatment of cancers, or the inhibition of tumor growth.
443111-01-19, 2,3-Bis(thiophen-2-yl)]quinoxaline-6-cerboxylic acid
[3-(morpholin-4-yl)propyl)amide 443111-36-29,
2,3-Bis(thiophen-2-yl)quinoxaline-6-cerboxylic acid [2-(chlorophenyl)ethyl]amide 443111-45-39, 2,3-Bis(thiophen-2-yl)quinoxaline-6-cerboxylic acid [2-(2-chlorophenyl)ethyl]amide 832080-84-59, 2-Chloro-1-(thiophen-2-yl)-6-quinoxalinecarboxylic acid [2-(2-chlorophenyl)ethyl]amide 832080-86-50-9,
2-[[2-(pyridin-4-yl)ethyl]amide 832080-86-19, 2-(gensyl)(methyl)aminol-3-(thiophen-2-yl)quinoxaline-6-carboxylic acid [2-(2-chlorophenyl)ethyl]amide 832080-06-61-9,
2-3-Bis(thiophen-2-yl)quinoxaline-6-carboxylic acid [2-(2-chlorophenyl)ethyl]amide 832082-01-59, 2,3-Bis(thiophen-2-yl)quinoxaline-6-carboxylic acid [2-(2-pyridyl)ethyl]amide 832082-01-59, 2,3-Bis(thiophen-2-yl)quinoxaline-6-carboxylic acid [2-(2-pyridyl)ethyl]amide 832082-01-59, 2,3-Bis(thiophen-2-yl)quinoxaline-6-carboxylic acid [2-(3-pyridyl)ethyl]amide 832082-03-09, 2,3-Bis(thiophen-2-yl)quinoxaline-6-carboxylic acid [2-(4-pyridyl)ethyl]amide 832082-03-09, 2,3-Bis(thiophen-2-yl)quinoxaline-6-carboxylic acid R-[3-(phenyl)propyl)amide RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapoutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of quinoxaline and pyrido[2,3-b]pyrazine derive. as PKB inhibitors for treatment of canceral

(Uses)
(preparation of quinoxaline and pyrido[2,3-b]pyrazine derive. as PKB inhibitors for treatment of cancers)
43111-01-1 CAPLUS
6-Quinoxalinecarboxamide, N-[3-(4-morpholinyl)propyl]-2,3-di-2-thienyl-(9C1) (CA INDEX NAME)

443111-36-2 CAPLUS 6-Quinoxalinecarboxamide, N-{2-(2-chlorophenyl)ethyl}-2,3-di-2-thienyl-(9CI) (CA INDEX NAME)

443111-45-3 CAPLUS 6-Quinoxalinecarboxamide, N-{2-(4-morpholinyl)ethyl}-2,3-di-2-thienyl-[9C1] (CA INDEX NAME)

832082-01-6 CAPLUS 6-Quinoxalinecarboxamide, N-{2-{2-methylphenyl}ethyl}-2,3-di-2-thienyl-[9C1] (CA INDEX NAME)

832082-02-7 CAPLUS 6-Quinoxalinecarboxamide, N-{2-(2-pyridinyl)ethyl}-2,3-di-2-thienyl- (9CI) (CA INDEX NAME)

832082-03-8 CAPLUS 6-Quinoxalinecarboxamide, N-[2-(3-pyridinyl)ethyl]-2,3-di-2-thienyl- (9CI) (CA INDEX RAME)

832082-04-9 CAPLUS 6-Quinoxalinecarboxamido, N-{2-(4-pyridinyl)ethyl}-2,3-di-2-thienyl- (9CI) (CA INDEX NAME)

832082-05-0 CAPLUS 6-Quinoxalinecarboxamide, N-(3-phenylpropyl)-2,3-di-2-thienyl- (9CI) (CA INDEX NAME)

832080-84-9 CAPLUS 6-Quinoxalinecarboxamide, 2-chloro-N-[2-(2-chloropheny1)ethy1]-3-(2-thieny1)- (9CI) (CA INDEX NAME)

832080-85-0 CAPLUS 6-Quinoxalinecarboxamide, N-[2-(2-chlorophenyl)ethyl]-2-[[2-(4-pyridinyl)ethyl]amino]-3-(2-thienyl)- (9CI) (CA INDEX NAME)

832080-86-1 CAPLUS 6-Quinoxalinecarboxamide, N-[2-(2-chlorophenyl)ethyl]-2-[methyl(phenylmethyl)amino]-3-(2-thienyl)- (9CI) (CA INDEX NAME)

832082-00-5 CAPLUS 6-Quinoxalinecarboxamide, N-{2-(2-methoxyphenyl)ethyl]-2,3-di-2-thienyl-(9CI) (CA INDEX NAME)

LI3 ANSWER 22 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2005:11522 CAPLUS
1005:11522 CAPLUS
112:261107
Preparation and redox properties of
N.N.N-1,3,5-trialkylated flavin derivatives and their
activity as redox catelysts
Linden, Auri A.; Hermanns, Nine; Ott, Sascha; Krueger,
Lare; Backvall, Jan-E.
CORPORATE SOURCE: Department of Organic Chemistry, Stockholm University,
Stockholm, 106 91, Swed.
Chemistry--A European Journal (2005), Volume Date
2004, 11(1), 112-19
CODEN: CEUJED; ISSN: 0947-6539
Miley-CCH Verlag GmbH & Co. KGAA
DOCUMENT TYPE: Journal
LANGUAGE: Endlish

PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): English CASREACT 147:261107

Inglish
RS SOURCE(S): English
RS SOURCE(S): CARRENCT 142:261107

Bight different flavin derivs. have been synthesized and the electronic effects of substituents in various positions on the flavin redox chemical were investigated. The redox potentials of the flavins, determined by cyclic voltementry, correlated with their efficiency as catalysts in the H202 oxidation of Me p-tolyl sulfide. Introduction of electron-withdrawing groups increased the stability of the reduced catelyst precursor.
843753-36-89 845753-39-19 845753-34-89

RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation and redox properties of N.N.N-1.3,5-trialkylated flavin derive. and activity as redox catelysts)
845753-36-8 CAPLUS
Benzo(g)peridine-7-carboxylic acid, 1,2,3,4-tetrahydro-1,3-dimethyl-2,4-dioxo-, butyl ester (SCI) (CA INDEX NAME)

Benzo[g]pteridine-7-carboxylic acid, 1,2,3,4-tetrahydro-2,4-dioxo-, butyl ester (SCI) (CA INDEX NAME)

845753-44-8 CAPLUS
Benzo[g]pteridine-8-carboxylic acid, 1,2,3,4-tetrahydro-2,4-dioxo-, butyl
ester (9CI) (CA INDEX NAMS)

845753-46-0 CAPLUS Benzo[g]pteridine-8-carboxylic acid, 1,2,3,4-tetrahydro-1,3-dimethyl-2,4-dioxor, butyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT:

45 THERE ARE 45 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSMER 23 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:1154708 CAPLUS
DOCUMENT NUMBER: 142:93840 of pyrido[1,2-e]pyrimidin-4-ones as
anticancer agents
INVENTOR(S): Wang, Weibo; Constantine, Ryan N.; Lagniton, Liena M.;

25 µM. The compns. that include a pharmaceutically acceptable carrier and one or more of the pyrido[1,2-a]pyrimidinyl compds. I, either alone or in combination with at least one addnl. therapeutic agent, were disclosed. 817205-84-89 RL: PRC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) [preparation of pyrido[1,2-a]pyrimidin-4-ones as anticancer agents) 817205-84-8 CAPUIS 6-Quinoxalinecarboxamide, N-(3-aminopropyl)-N-[1-[4-oxo-3-(phenylmethyl)-4H-pyrido[1,2-a]pyrimidin-2-yl]propyl]- (9CI) (CA INDEX NAME)

(CH2) 3 - NH2

L13 ANSWER 24 OP 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
1004:976488 CAPLUS
1171T.R:
142:130137
15TH.R:

s24960-70-5 RL: ARC (Analytical reagent use); ANST (Analytical study); USES (Uses) (serum 3-deoxyglucosome is higher in streptozotocin-induced diabetic

rate)
84-90-70-5 CAPLUS
64-90inoxalinecarboxamide, N-[4-ox0-4-[[2-(2-pyridinyldithio)ethyl]amino]bu
tyl]-3-(2.3,4-trihydroxybutyl)- (9CI) (CA INDEX NAME)

DOCLOGNT TYPE:

Pecchi, Sabina; Burger, Matthew T.; Desai, Manoj C. Chiron Corporation, USA PCT Int. Appl., 78 pp. CODEN: PIXLO2

PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | | TENT | | | | | | | | | | | | | | | | |
|------|------|-------|------|-----|-----|-----|-----|------|------|-----|------|-------|-------|-----|-----|-----|------|-----|
| | | | | | | | | | | | | | | | | | | |
| | WO | 2004 | 1133 | 35 | | A2 | | 2004 | 1229 | 1 | WO 2 | 004- | US19 | 158 | | 2 | 0040 | 617 |
| | MO | 2004 | 1133 | 35 | | A3 | | 2005 | 0324 | | | | | | | | | |
| | | W: | AE, | AG, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BR, | BW, | BY, | BZ, | CA, | CH, |
| | | | CN, | œ, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FI, | GB, | GD, |
| | | | GE. | GH. | GM. | KR. | HU. | ID. | IL. | IN. | IS. | JP. | KE. | KG. | KP. | KR. | KZ. | LC. |
| | | | LK. | LR. | LS. | LT. | LU. | LV. | MA. | MD. | MG. | MK, | MN. | MW. | MX. | MZ. | NA. | NI. |
| | | | | | | | | | | | | BC. | | | | | | |
| | | | | | | | | | | | | UZ. | | | | | | |
| | | RW: | | | | | | | | | | SL, | | | | | | |
| | | | | | | | | | | | | BE, | | | | | | |
| | | | | | | | | | | | | LU, | | | | | | |
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| | | | | TD. | | | | ٠., | , | , | , | | ***, | , | •, | , | , | |
| | AU | 2004 | | | | | | 2004 | 1229 | - 1 | AU 2 | 004- | 2497 | 30 | | 24 | 0040 | 617 |
| | | 2528 | | | | | | | | | | | | | | | | |
| | US | 2005 | 0854 | 90 | | A1 | | 2005 | 0421 | | US 2 | 004- | 87070 | 7 | | 20 | 0040 | 617 |
| | | 1636 | | | | | | | | | | | | | | | | |
| | | R: | AT. | BE. | CH. | DE. | DK. | ES. | FR. | GB. | GR. | IT. | LI. | LU. | NL. | SE. | MC. | PT. |
| | | | | | | | | | | | | KU, | | | | | | |
| PRIO | DIT | Y APP | | | | | | | , | | | 003- | | | | 2 2 | 0030 | 620 |
| | | | | | | | | | | | | 004-1 | | | | | | |
| | R SC | | | | | | | | | | | | | | | | | |

The title compds. I [R1 = H, alkyl, aryl, etc.; R2, R3 = H, alkyl, aryl, etc.; or R2 and R3 taken together with the carbon atom to which they are attached form a 3-7 membered carbocyclic or heterocyclic ring; R4 = H, alkyl, aryl, etc.; R5 = H, alkyl, aryl, etc.; R6-R9 = H, halo, N02, etc.], useful, either alone or in combination with at least one addnl. therapeutic agent, in the prophylaxis or treatment of proliferative diseases, were prepared E.g., a multi-step synthesis of II, starting from 2-aminopyridine and Et 4-chloroacetoacetate, was given. Certain compds. I were shown to have a KSP inhibitory activity at an IC50 of less than about

PAGE 1-B

824960-65-8 824960-66-9 824960-67-0
824960-68-1 823637-48-7
RL: BBU (Biological study, unclassified); PRP (Properties); BIOL
(Biological study)
(serum 3-deoxyglucosone is higher in streptoxotocin-induced diabetic rate)
824960-65-8 CAPLUS
L-Lysine, N2-(6-[[5-[(3s.48,6R)-hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl]-10-xopentyl]amino]-1-oxohexyl]-N6-[[2-(2,3,4-trihydroxybutyl)-6-quinoxalinyl]carbonyl]- (9CI) (CA INDEX NAME)

PAGE 1-B

824960-66-9 CAPLUS
L-Lysine, N2-[6-[[5-[(3aS,48,6aR)-hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl]-l-oxopentyl]amino]-1-oxohexyl]-N6-(6-quinoxalinylcarbonyl)- (9CI)
(CA INDEX NAME)

PAGE 1-A

PAGE 1-B

824960-67-0 CAPLUS
L-Lysine, N2-[6-[[5-[(385,48,68R)-hexahydro-2-oxo-1H-thieno{3,4-d]imidazol-4-yl]-1-oxopenty]|amino]-1-oxohexyl]-N6-{(2-methyl-6-quinoxalinyl)carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

824960-68-1 CAPLUS
L-Lysine, M6-{[2-(2,3-dihydroxypropyl)-6-quinoxalinyl]carbonyl]-N2-{6-{[5-(36,46,66R)-hexahydro-2-oxo-1H-thieno[3,4-d]imidezol-4-yl]-1-oxopentyl]amino]-1-oxohexyl]- (9CI) (CA INDEX NAME)

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 25 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
111:395806 Preparation of quinoxalinyl macrocyclic hepatitis C
esrine protease inhibitors
Nakajima, Suanne; Sun, Ying; Tang, Datong; Xu, Gouyou;
PATENT ASSIGNES(S):
BATENT ASSIGNES(S):
SOURCE:
PATENT ASSIGNES(S):
COCEN: PIXXD2
DOCUMENT TYPE:

L13 ANSWER 25 OF 181
CAPLUS COPYRIGHT 2006 ACS on STN
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COCEN: PIXXD2
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COCEN: PIXXD2
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ACCENTING TO THE ACCENT ASSIGNES ACCENT

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | PA | TKAT | NO. | | | KIN | D | DATE | | | APPL | ICAT | ION | NO. | | D. | ATE | | |
|---|-----|------|------|------|-----|-----|-----|------|------|-----|-------|------|------|-----|-----|-----|------|------|---|
| | | | | | | | - | | | | | | | | | - | | | |
| | WO | 2004 | 0937 | 98 | | A2 | | 2004 | 1104 | | WO 2 | 004- | US11 | 841 | | 2 | 0040 | 116 | |
| | WO | 2004 | 0937 | 98 | | A3 | | 2005 | 1208 | | | | | | | | | | |
| | | W: | AE, | AG, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BR, | BW. | BY, | BZ, | CA. | CH, | |
| | | | CN, | co. | CR. | CU. | CZ. | DE, | DK, | DM, | DZ. | EC. | EE. | EG. | ES. | PI. | GB. | GD, | |
| | | | | | | | | ID. | | | | | | | | | | | |
| | | | LK. | LR. | LS. | LT. | LU. | LV. | MA. | MD. | MG. | MK. | MOV. | MW. | MX. | MZ. | NA. | NI. | |
| | | | NO. | NZ. | OM. | PG. | PH. | PL, | PT. | RO. | RU. | SC. | SD. | SE. | SG. | SK. | SL. | SY. | |
| | | | | | | | | TZ, | | | | | | | | | | | |
| | | RW: | | | | | | MW, | | | | | | | | | | | |
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| | MA | 2004 | | | | 8.1 | | 2004 | 1104 | | AII 2 | 004- | 2319 | . 7 | | 2 | 0040 | 116 | |
| | | 2522 | | | | | | | | | | | | | | | | | |
| | | | | | | | | | | | | | | | | | | | |
| | | 2004 | | | | | | | | | | | | | | | | | |
| | EP | 1615 | 613 | | | A2 | | 3006 | 0118 | | BP 2 | 004- | 7502 | 36 | | 2 | 3040 | 116 | |
| | | R: | AT, | BE, | CH, | DE. | DK, | BS, | FR, | GB, | GR, | IT. | LI, | LU, | NL, | SE. | MC, | PT, | |
| | | | IE. | SI. | LT. | LV. | FI. | RO. | MK. | CY. | AL. | TR. | BG. | CZ. | EE. | HU. | PL. | SK. | H |
| c | RIT | APP | LN. | INFO | | | | | | - 1 | US 2 | 003- | 4187 | 59 | | A 2 | 0030 | 116 | |
| | | | | | | | | | | | | | | | | | | | |

OTHER SOURCE(S): MARPAT 141:395806

The invention relates to macrocyclic compds. I [A is H, CO2R1, COR2,

Absolute stereochemistry.

PAGE 1-B

825637-48-7 CAPLUS L-Lysine, N2-[6-[[5-[(38,48,68R)-hexahydro-2-oxo-1H-thieno(3,4-d]imidazol-4-yl]-1-oxopenty]|amino]-1-oxohexyl]-N6-[[2-[(1R,28,3R]-1,2,3,4-tetrahydroxybutyl]-6-quinoxalinyl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS

COMMR2. CSNMR2 or SO2R2; G is CH. alkoxy. NHSO2R1. COR2. CO2R1 or COMMR2; L is S. SCN2. SO2. O. COCM2. CHMeCN2 stc.: m. n = 0-2; p = 0-4; R2 is a bond or H2; R1 is H. (un) substituted alkyl. alkenyl. alkynyl. cycloslkyl. aryl. arylakyl. heteroaryl. heteroarylakyl or heterocycloslkyl; R2 is any group given for R1 or mono- or dielkylamino or -arylamino; R3. R4 not defined; X and Y taken together with the carbon atoms to which they are attached form (un) substituted aryl or heteroaryl; M is absent, O. S. NH, C(O)NR1 or NR1; Z is H, -CN, -SN, -NSO, -NSS, MHHR2, N3, halo, cycloslkyl, aryl, etc.] or their pharmaceutically-acceptable salts, esters or prodrugs which inhibit serine protesse activity, particularly the activity of hepatitis C virus (HCV) NS3-NS4A protease. The compds. of the invention interfere with the life cycle of the hepatitis C virus and are also useful as antiviral agents. Thus, macrocycle II (Boc = tert-butoxycarbonyl) was prepared via peptide coupling and ring-closing metathesis reactions.

197600-64-0p
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); SIOL (Biological study); PREP (Preparation); USES (Uses) (Uses) (Preparation of quinoxalinyl cyclic peptides as hepatitis C serine protease inhibitors)

107600-64-0 CAPLUS
(Vologropa (e) pyrrolo[1,2-a][1,4]diazacyclopentadecine-14a(SH)-carboxylic acid. 6-[([1,1-dimethylethoxylearbonyl]amino].

1,2,3,6,7,8,9,10,11,11a,14,15,16,16a-tetradechydro-5,16-dioxo-2-[[6-[([henylesthyl)amino]carbonyl]-1-(2-thienyl)-2-quinoxalinyl)oxyl-, (2R,6S,11aS,14aR,16aS)-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

PAGE 1-A

PAGE 2-B

L13 ANSWER 26 OP 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:857399 CAPLUS
DOCUMENT NUMBER: 141:343478
Use of small molecule compounds for immunopotentiation
Valiante, Nicholaes
Chiron Corporation, USA
PATENT ASSIGNEE(S): PTXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUMBURACION: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT | NO. | | | KIN | ם | DATE | | | APPL | ICAT | ION | NO. | | D. | ATE | |
|---------|------|-----|-----|-----|-----|------|------|-----|------|------|------|-----|-----|-----|------|-----|
| | | | | | - | | | | | | | | | - | | |
| WO 2004 | 0871 | 53 | | A2 | | 2004 | 1014 | | WO 2 | 004- | US10 | 331 | | 2 | 0040 | 329 |
| WO 2004 | 0871 | 53 | | A3 | | 2005 | 0317 | | | | | | | | | |
| W: | AE, | AG, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BR, | BW, | BY, | BZ, | CA, | CH. |
| | CN, | co, | CR, | Cυ, | CZ. | DE, | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FI, | GB, | GD, |
| | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | is, | JP, | KE, | KG, | KP, | KR, | KZ, | LC, |
| | LK, | LR, | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NA, | NI, |
| | NO, | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | BK, | SL, | SY, |
| | TJ, | TM, | TN. | TR, | TT, | TZ, | UA, | υσ, | US, | υz, | VC, | VN, | YU, | ZA, | ZM, | ZW |
| RW: | BW, | GH, | GM, | KB, | LS, | MW, | MZ, | SD, | SL, | SZ. | TZ, | UG, | ZM, | ZW, | AM, | AZ, |
| | BY, | KG, | KZ, | MD, | RU, | TJ, | TM, | AT, | BE, | BG, | CH, | CY, | CZ, | DE, | DK, | EE, |
| | ES, | FI, | FR, | GB, | GR, | HU, | IB, | IT, | LU, | MC, | ΝL, | PL, | PT, | RO, | SE, | SI, |

M.; Osgood, Stephen A.; Bertrand, Myra; Swayze, Eric

CORPORATE SOURCE:

SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S):

M.; Osgood, Stephen A.; Bettrand, Myra; Swayze, Eric S.

PORATE SOURCE: Ibis Therapeutics, Isis Pharmaceuticals, Inc., Carlebad, C. A. 92008. USA

RCE: Bicorranic & Medicinal Chemistry Letters (2004), 14(21), 5257-5261

CODEN: BMCLE8; ISEN: 0960-894X

LISHER: Bleevier B.V.

LUMENT TYPE: Journal

UNIONE TYPE: Journal

RR SOURCE(S): CASREACT 142:3310

We report on lead optimization of a compound that was originally discovered to bind bacterial 235 RRNA near the L11 binding site and inhibit translation in vitro, but lacked detectable antibacterial activity. In this study, we were able to generate compds, with antibacterial activity against Gran-neg, and Gran-pos, pathogens, including a methicillin-resistant Staphylococcus aureus strain. 737770-86-49

RL: BSU (Biological study, unclessified); SPN (Synthetic preparation); BIDLO (Biological study); PREP (Preparation)

(optimizing the antibacterial activity of a lead structure)
797770-86-5 CAPLUS

6-Quinovalinesarboxanids, N-[1-[4-[3]-(dimethylamino) propoxy) phenyl]-2-oxo-

/s://u-ss-s CAPLUS
6-Quinoxalinacarboxasida, N-[1-[4-[3-(dimethylamino)propoxy]phanyl]-2-oxo-2-(1-piperazinyl)ethyl]-1,2,3,4-tetrahydro-2,3-dioxo-(SCI) (CA INDEX NAMS)

THERE ARE 17 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 28 OF 181 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: CAPLUS COPYRIGHT 2006 ACS ON STN 2004:515470 CAPLUS 141:71352

141:71352
Preparation of biphenylaminobenzoates and related compounds as modulators of peroxisome proliferator activated receptor y (PPARy) type receptors as drugs and cosmetics.
Clary, Leurence; Collette, Pascal; Rivier, Michel; Jomard, Andre Calderma Research & Development, S.N.C., Fr. PCT Int. Appl., 50 pp.
CODEN: PIXXO2
Patent English
2

INVENTOR (S):

PATENT ASSIGNEE (S) : SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. CO PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

MO 2004052840 A1 20040624 NO 2003-EP15010 20031211

M: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DR, DM, DM, DZ, EC, EE, EG, EB, FI, GB, GD, GR, GH, GM, HR, HU, ID, IL, IN, IS, JP, KR, KG, KP, KR, KZ, LC,

SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GO, GM, ML, MR, NE, SN, TD, TG

CA 2520124 AA 20041014 CA 2004-2520124 20040329

US 2005136065 A1 20050621 US 2004-814480 20040329

FF 1608169 A2 20051226 BP 2004-758593 20040329

R: AT, BE, CH, DB, DK, ES, FR, GB, GR, IT, LI, LU, NL, EE, MC, PT, IE, SIL, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, RU, PL, SK

PRIORITY APPLN. INFO.: US 2001-458888P P 20030328

WO 2004-US10331 OTHER SOURCE(S): MARPAT 141:343476

The invention provides immunostimulatory compns. comprising a small mol. immunopotentiator (SMIP) compound and methods of administration thereof. Also provided are methods of administering a SMIP compound in an effective amount to enhance the immune response of a subject to an antigen. Purther provided are compns. and methods of administering SMIP compds. alone or in combination with another sgent for the treatment of cancer, infectious diseases and/or allergies/asthms. Preparation of selected compds., e.g. I, is included. 654634-24-9

KL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USSS (Uses)

(small mol. compds. for immunopotentiation)
654634-24-9 CAPLUS
Benzo(b)phenazine-2-carboxylic acid. 6.11-41bd-2-6-11-41cc.

Benzo[b] phenazine-2-carboxylic acid, 6,11-dihydro-6,11-dioxo-, ethyl ester (9CI) (CA INDEX NAME)

L13 ANSWER 27 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:791916 CAPLUS

142:3310

DOCUMENT NUMBER: TITLE: 142:33:0
Optimizing the antibacterial activity of a lead
structure discovered by SAR by MS technology
Jefferson. Elizabeth A.; Seth. Punit P.; Robinson,
Dale E.; Winter, Dana K.; Miyaji, Alycia; Risen, Lisa AUTHOR (S)

LK, LR, LS, LT, LU, LV, MA, MD, MG, MX, MN, MM, MX, MZ, NI, NO, NZ, OM, MO, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TO, TM, TM, TM, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZM, RM, BM, GM, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, ZM, ZM, AM, AZ, BY, KG, KZ, MG, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, NG, EE, SF, FF, GB, GR, HU, IE, IT, LU, MC, MC, PT, RO, SE, SI, SK, TG, TF, RF, BF, BJ, CF, CG, CI, CM, QA, OM, QG, OM, ML, MR, MR, SK, TD, TG, AZ, SCO6732 AA 20040624 CA 2003-2505732 200312311 CA 2506732 AA 20040624 CA 2003-2506732 200312311 RF, ST, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, ES, MU, SK, BR 200316193 A 20050927 BR 2003-261593 200312311 JP, 2006509798 T2 20050231 JP, 2004-550699 200312311 JP, 2006509798 T2 200600944 A1 20060012 BR 2003-161593 200312311 PRIORITY APPLIN. INFO: FR 2006115 ST 200601215 ST 200601215 ST 2006001215 ST 2006001215 ST 200601215 ST 200601215 ST 200601215 ST 200601215 ST 200601215 ST 2006012121 ST 200601215 S

MARPAT 141:71352

OTHER SOURCE(S):

ies] (claimed compound; preparation of biphenylaminobenzoates and related compds. as modulators of peroxisome proliferator activated receptor γ)

$$\begin{array}{c|c} & \stackrel{\text{Me } 0}{\text{C}} \\ & \stackrel{\text{NH}}{\longrightarrow} \\ & \stackrel{\text{CN}_2}{\longrightarrow} \\ & \stackrel{\text{NH}}{\longrightarrow} \\ & \stackrel{\text{NH}}{\longrightarrow}$$

706779-85-3 CAPLUS
Senzoic acid, 3-[[3'-[[methyl(6-quinoxalinylcarbonyl)amino]methyl][1,1'-biphenyl]-4-yl]amino]-, 2-methylpropyl ester (9CI) (CA INDEX NAME)

706779-92-2 CAPLUS
BenZoic acid, 3-{[3'-{[methyl(6-quinoxalinylcarbonyl)amino]methyl][1,1'-biphenyl]-4-yl]oxy]-, 2-{4-morpholinyl)ethyl eater (9CI) (CA INDEX NAME)

L13 ANSMER 29 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:493124 CAPLUS
DOCUMENT NUMBER: 141:59705
TITLB: New amino biphenyl compounds as modulators of peroxiscome proliferator-activated y-receptors
(PPAR y) for commettio or pharmacoutical

(PPAR y) for commette of pursuance compositions
Clary, Laurence; Collette, Paacal; Rivier, Michel;
Jomard, Andre
Galderma Research & Development, Fr.
Pr. Demande, 47 pp.

INVENTOR (S):

PATENT ASSIGNEE (S):

CODEN: FRXXBL Patent French DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

DATE

2001212
20031211
20031211
BY, BZ, CA, CH,
BS, FI, GB, GD,
KP, KR, KZ, LC,
KX, MZ, MI, NO,
SK, SL, SY, TJ,
ZA, ZM, ZM,
ZM, ZM, AM, AZ,
CZ, DE, DK, EE,
RO, SE, SI, SK,
MR, NE, SN, TD, TO
20031211
NL, SE, MC, PT, BR 2003016193 JP 2006509798 US 2006009484 PRIORITY APPLN. INFO.:

IN SOURCE(8): MARRAT 141:59705 W0 2003-EP15010 W 20031211

Nevamino biphenyl compds a PPAR y receptor modulatora are prepared of a deficiency for compact compact and product compact OTHER SOURCE(S):

706779-85-3 CAPLUS
Benzoic acid, 3-[(3'-[[methyl(6-quinoxalinylcarbonyl)amino]methyl][1,1'-biphenyl]-4-yl]amino]-, 2-methylpropyl ester (9CI) (CA IMDEX NAME)

706779-92-2 CAPLUS

Benzoic acid, 3-[[3'-[[methyl (6-quinoxalinylcarbonyl)amino]methyl][1,1'-biphnyl]-(-4-yl)oxyl-, 2-(4-morpholinyl)ethyl ester (9CI) (CA INDEX NAMS)

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THI RECORD. ALL CITATIONS AVAILABLE IN THE RE FORM.

ACCESSION NUMBER: 1004:467877 CAPLUS 1004:467877 CAPLUS 141:34617 TITLE: Preparation of Diphenyla which activate peroxisome proliferator activated receptor- y (PPARy) receptora for use in druga and coametics. Calley, Laurence: Boulx-Pater, Claire; Rivier, Michel; Collette, Pascal; Jonard, Andre Galderma Research & Development, S.N.C., Fr. PCT Int. Appl., 73 pp. CODEN: PIXXOZ PATENT INFORMATION: 2

PATENT INFORMATION: 2

PATENT INFORMATION: PATENT NO.

ES, FI, PR, GB, GR, HU, IS, IT, LU, MC, NL, PT, RO, SE, SI, SK, TB, PR, 2647580 A1 2004052B FR 2002-14793 20021125

FR 2647580 B1 2004052B FR 2002-14793 20021125

CA 25056523 AA 20040610 CA 2003-2506523 20021121

AU 2003294025 A1 20040618 AU 2003-294025 20011121

AU 2003294025 A1 20040618 AU 2003-294025 20011121

FI 1567509 A2 20050931 FP 2003-794039 20011121

R: AT, BE, CH, DE, DK, SE, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IS, SI, LT, LV, FR, OMK, CY, AL, TR, BG, CZ, EE, HU, SK

BR 2003015951 A 20050013 BR 2003-15951 20051121

UZ 2006004048 A1 20060105 US 2005-135499 2005524

PRIORITY APPLN. INFO::

US 2002-14793 A 20021125 US 2002-430698P WO 2003-EP15002 OTHER SOURCE(S): MARPAT 141:38617

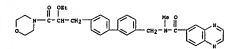
Title compds. [I; R1 = alkyl, (substituted) Ph, N-protected α-amino acid, etc.; R2 = Q1, Q2, COR8; R3 = H, halo, alkyl, OH, alkoxy, aralkoxy, aryloxy, polyether residue, NO2, amino; R4 = alkyl, aryl, aralkyl, heteroaryl, heterocyclyl, etc.; A = (CH2)x(NR12)y(CO)xDw, etc.; w, x, y, z = Q, 1; D = Q, S, G12, NR13; V = Q, N, S; M = N, CR10; Y = N, C, R7- H, alkyl, aryl, aralkyl, heteroaryl, heterocyclyl, 88 = Q(CH2)xR9. OH, alkoxy, aralkoxy, aryloxy, aryl, aralkyl, heteroaryl, heterocyclyl, etc.; R10 = H, alkyl, aryl, aralkyl, heteroaryl, heterocyclyl, etc.; R10 = H, alkyl, aryl, aralkyl, heteroaryl, heterocyclyl; R12, R13 = H, alkyl, R14 = halo; v = 1-31, were prepared Thua, HATU, P3-carbodiimide reain, (4--[2-ethoxy-2-(5-propyl-1,2,4-oxadiasol-2-yllethyl] biphenyl-3-ylmethylmethylamine (preparation given), and 6-(2-methoxyethoxymethoxy)naphthalene-2-carboxylic acid were stirred together in DMY/CH2C13 tog 18 h at room temperature to give a residue which was stirred 5 h with NP-carbonate reain in DMY/CH2C13 to give 100% N. (4-'[2-ethoxy-2-(5-propyl-1,2,4-oxadiasol-2-yl)ethyl)piphenyl-3-ylmethyl)-N-methyl-6-(2-methoxyethoxymethoxy)naphthalene-2-carboxamide. In a croasead-curve PPAR y activation test, the latter showed Kd app = 250 nM. 632780-24-2P AR

250 nM.

(\$2780-94-2P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Usea)

(claimed compound; preparation of biphenyls which activate peroxisome proliferator activated receptor- γ receptors for use in drugs and commetics)

(52780-94-2 CAPLUS
6-Quinoxalinecarboxamide, N-{{4'-{2-ethoxy-3-{4-morpholinyl}-1-oxopropyl}{1,1'-biphenyl}-3-yl}methyl}-N-methyl- (9CI) (CA INDEX NAME) IT



L13 ANSWER 31 OF 181
ACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE:

TITLE:

TITLE:

INVENTOR(8):

EARTHWT ACCIGNET(8):

PARTENT ACCIGNET(8):

L2004:433750 CAPLUS
CAPLU

PATENT ASSIGNEE (S) : SOURCE:

NEIMOTOR, DAVIC. USA U.S. Pat. Appl. Publ., 121 pp., which CODEN: USXXCO Patent English 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE 20031003 P 20021030 P 20030407 US 2003-678565 US 2002-422312P US 2003-460911P 20040527 A1 US 2004102360 PRIORITY APPLN, INFO.: MARPAT 141:7131

OTHER SOURCE(S):

STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The present invention relates to methods of treating cancer using a combination of at least two Akt inhibitors I (wherein O = (un) substituted heterocycly), aryl; U, V, W, and X = independently CH, N, Y, Z = independently CH, N, provided that at least one of Y and Z = N; n = 0-3; p = 0-2; q = 0-4; R1, R2, R7 = independently halo, CN, OR, CKO, NO2, or (un) substituted (cyclo) alkyl (oxy), alkenyl (oxy), alkynyl (oxy), alkyn

PAGE 1-B

CRN 76-05-1 CMF C2 H F3 O2

_612847-42-4 CAPLUS 6-Ouinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-3-[4-[[4-(2,3-dihydro-2-oxo-H+benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl- (9CI) (CA INDEX NAME)

612847-43-5 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-2-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-3-phenyl-(9CI)
(CA INDEX NAME)

612848-56-3 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-2-[4-[[4-(2,3-dihydro-2-oxo-1R-benzimidaxol-1-yl)-1-piperidinyl]eethyl]phenyl]-3-phenyl-, trifluoroacetate (9CI) (CA INDEX NAME)

CM 1

CRN 612847-43-5 CMF C40 H43 N7 O2

with chloroform and ethanolic HCl. III =HCl, a selective Akt1 and Akt2 inhibitor, demonstrated a 3.2-fold in caspase 3 activation over control compared to a 1.2-fold increase for a protein kinase inhibitor. Combination treatment produced a 9-fold increase in caspase 3 activation. 612847-33-39 612847-34-49 612847-42-49 612847-437-9 612848-63-9 612848-53-9 612848-51-9 foldes-5-39 foldes-5-

(Uses)

(antitumor agent; preparation of quinazolines and analogs as Akt inhibitors and indoles as protein kinase inhibitors for use in synergistic combination therapy for treatment of cancer)
612647-33-3 CAPLUS
6-Quinoxalinecarboxamide, 3-{4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-y])-1-piperidiny]|methyl]phenyl]-N-{3-(1H-imidazol-1-y])propyl]-2-phenyl(9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-E

612847-34-4 CAPLUS
6-Quinoxalinecarboxamide, 3-[4-[[4-[2,3-dihydro-2-oxo-1H-benzimidazol-1-yl]-1-piperidinyl]methyl]phenyl]-N-[3-(1H-imidazol-1-yl)propyl]-2-phenyl-, bis(trifiluoroacetate) (9C1) (CA INDEX NAMS)

CM 1

CRN 612847-33-3 CMF C40 H38 N8 O2

CM 2

CRN 76-05-1 CMF C2 H F3 O2

612848-57-4 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-3-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl]-1-piperidinyl]methyl]phenyl]-2-phenyl-,
trifluoroacetate (9CI) (CA INDEX NAME)

CRN 612847-42-4 CMF C40 H43 N7 O2

CH 2

CRN 76-05-1 CMF C2 H F3 O2

612848-59-6 CAPLUS
D-arabino-Hexose, 2-deoxy-2-[[[2-[4-[[4-(2,3-dihydro-2-oxo-1H-bensimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-3-phenyl-6-quinoxalinyl]carbonyl]amino]-, (2\(\xi\)-, trifluoroacotate (salt) (9CI) (CA INDEX NAME)

CH 1

CRN 612848-58-5 CMF C40 H40 N6 O7

Absolute stereochemistry.

PAGE 1-B

CRN 76-05-1 CMF C2 H F3 O2

612848-61-0 CAPLUS
D-arabino-Hexose, 2-deoxy-2-[{3-[4-[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-y1)-1-piperidinyl]methyl]phnyl]-2-phenyl-6-quinoxalinyl]carbonyl]mino]-, (2\$)-, trifluoroacetate (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 612848-60-9 CMF C40 H40 N6 O7

Absolute stereochemistry.

L13 ANSWER 32 OF 181 CAPLUS COPYRIGHT 3006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
140:432795 CAPLUS
TITUE:
PATENT ASSIGNER(S):
PATENT ASSIGNER(S):
COLLETT, PACENT ASSIGNER(S):
DOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
DATENT ASSIGNER(S):
PATENT ASSIGNER(S)

LANGUAGE: FAMILY ACC. NUM. COUNT:

| | | | | | | | | | | | LICAT | | | | | ATE | | |
|-----|------|------|------|-----|-----|-----|------|------|-----|----|----------------|------|-----|-----|------|-------|-----|--|
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| | 2847 | | | | | | | | | FR | 2002- | 1479 | 3 | | 2 | 0021 | 125 | |
| | 2847 | | | | | | | | | | | | | | | | | |
| CA | 2506 | 523 | | | AA | | 2004 | 0610 | | CA | 2003- | 2506 | 523 | | 2 | 3031 | 121 | |
| | | | | | | | | | | WO | 2003- | EP15 | 002 | | 21 | 3031 | 121 | |
| WO | 2004 | 0483 | 51 | | A3 | | 2004 | 0812 | | | | | | | | | | |
| | ₩: | | | | | | | | | | , BG, | | | | | | | |
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| | RW: | | | | | | | | | | , sz, | | | | | | | |
| | | | | | | | | | | | , BG, | | | | | | | |
| | | | | | | | | | | | , MC, | | | | | | | |
| | | | | | | | | | | | , GQ, | | | | | | | |
| | | | | | | | | | | | 2003- | | | | | | | |
| ₽P | | | | | | | | | | | 2003- | | | | | | | |
| | R: | | | | | | | | | | . IT. | | | | | | | |
| | | IB. | BI, | LT. | LV, | PI. | RO, | MK, | CY, | AL | , TR, | BG, | cz, | EE, | HU, | sĸ | | |
| BR | 2003 | 0159 | 51 | | A | | 2005 | 0913 | | BR | 2003- | 1595 | 1 | | 2 | 3031 | 121 | |
| US | 2006 | 0040 | 48 | | A1 | | 2006 | 0105 | - | US | 2005-
2002- | 1354 | 99 | | 21 | 00509 | 524 | |
| ITY | APP | LN. | INFO | . : | | | | | | FR | 2002- | 1479 | 3 | - 1 | A 21 | 3021 | 125 | |
| | | | | | | | | | | | 2002- | | | | | | | |

PAGE 1-B

CM 2

CRN 76-05-1 CMP C2 H F3 O2

695816-06-9 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-2-[4-[(4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-N-methyl-3-phenyl-trifluoroacetate (9CI) (CA INDEX NAME)

CM 1

CRN 695816-05-8 CMF C41 H45 N7 O2

CM 2

The invention relates to new biarom. compds. I and their method of preparation, and their use in cosmetic or pharmaceutical compns. intended for use in human or veterinary medicine (such as cardiovascular diseases, immunity diseases and/or diseases related to the metabolism of the lipids). Thus, N-(4--(2-ethoxy-2-(5-propy)-1-(1,3,4)oxadiarol-2-y-1)-ethyl]-biphenyl-3-dimethyl]-N-methyl-4-phenoxy-benzoine (sid with (4--(2-ethoxy-2-(5-propy)-5 [1,3,4]oxadiarol-2-y-1)-methyl]-methyl-amine. The specific affinity of the composition for PPAR- y is shown. A tablet contained II 0.001, starch 0.114, disalcium phosphate 0.020, silica 0.020, lactose 0.030, tale 0.010, and magnesium stearete 0.005 g. S2780-342. All of the composition for PPAR- y is shown. A tablet contained II 0.001, starch 0.114, disalcium phosphate 0.020, silica 0.020, lactose 0.030, tale 0.010, and magnesium stearete 0.005 g. S2780-3-2 RL: COS (Commetic use): PAC (Pharmacological activity); THU (Therapeutic use): BIOL (Biological study): USES (Uses) (preparation of biaros. compds. as activators of PPAR receptors and their uses in commetic or pharmaceutical compns.)
S92780-34-2 CAPIUS
6-Quinoxalinecarboxanide, N-[{4-[2-ethoxy-3-(4-morpholinyl)-3-oxopropyl][1,1'-biphenyl]-3-yl]methyl]-N-methyl- (SCI) (CA INDSX NAME)

REFERENCE COUNT: THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMA

ACCESSION NUMBER:
DOCUMENT NUMBER:
1TITLE:
1NVENTOR(S):
1A:7139
Preparation of indolylquinoxalinones for treating hyperproliferative disorders and diseases associated with amjogenesis
Ladouceur, Gastan H.; Bear, Brian; Bi, Cheng; Brittelli, David R.; Burke, Michael J.; Chen, Gang; Cook, James; Dumas, Jacques; Sibley, Robert; Turner, Michael R.

BAYENT ASSIGNEE(S):
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
PARILY ACC. NUM. COUNT:
1

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

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PATENT NO.
                                                                                                                                                                                                                                                                                  DATE
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                                       DATE APPLICATION NO. DATE DATE APPLICATION NO. DATE DATE

MO 2000033950 A1 20040527 MO 2003-U338003 20031110

M: AS. AG. AL. AM. AT. AU. AZ. BA. BB. BG. BR. BY. BZ. CA. CH. CN. CO. CR. CU. CZ. DE. DK. DM. DZ. EC. EE. BG. EB. FI. GB. GD. GE. GE. GG. GH. GM. HER. HU, ID. IL. IN. II. II. JP. KE. NG. KP. KR. KZ. LC. LK. LR. LS. LT. LU, LV. MA. MD. MG. MK, MN. MM. MK. MZ. NK. KZ. LC. LK. TN. TN. TR. TT. TZ. UA. UG. US. UZ. VC. VN. VY. UZA. ZM. ZM. TN. TN. TR. TT. TZ. UA. UG. US. UZ. VC. VN. VY. UZA. ZM. ZM. AZ. BY. KG. KZ. NG. RU. TJ. TM. AT. BE. BG. CH. CY. CZ. DE. DK. BE. ES. MR. KY. DE. MY. CA. CS. DE. CS. TY. UG. ZM. ZM. AM. AZ. ES. TR. FF. BJ. CF. CG. CI. CM. GA. GN. GQ. GM. ML. KR. NR. SN. TD. TG. CA. 2505819 AA 2004527 CA. 2003-2505419 20031110

AU 2002320744 A1 20046633 AV 2003-250544 20031110

AU 2002320744 A1 2004663 AV 2003-250544 20031110

R: AT. BE. CH. DE. DK. ES. FR. GB. GR. IT. LI. LU, ML. RS. SN. MC. PT. ER. AT. BE. LL. CV. VY. FR. AG. KC. CY. AL. TR. BG. CZ. EE. HU, SK. CY. AL. TR. BG. CG. CS. EE. HU, SK. CY. AL. TR. BG. CG. CS. EE. HU, SK. CY. AL. TR. BG. CG. CS. EE. CO031110

AD 2006509401 A1 20065023 DY 2005-557146 20031110

AD 2006509401 A1 20065025 US 2005-557146 20031110
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BR 2003-16169
CN 2003-30108639
JP 2005-507146
US 2005-534215
NO 2005-2796
US 2002-425490P
US 2003-460915P
US 2003-464202P
MO 2003-US36003
BR 2003016169
CN 1738814
JP 2006509840
US 2006004011
NO 2005002796
PRIORITY APPLN, INPO.:
                                                                                                                                                                                                                                                                                    20060105
20050609
OTHER SOURCE(S):
                                                                                                                                                                                                                     MARPAT 141:7139
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The invention relates to title compds. I [wherein Ar = 6-membered eromatic ring containing 0-2 N atoms; R1 and R2 = independently H, halo, CF3, acyl, piperidinyl, piperazinyl, morpholinyl, or (un)substituted elkyl, alkoxy, amino, pyrrolidinyl, Ph, etc.; R3 = H, elkyl, OH, NOZ, NR3, elkylemino, alkoxyamino, or (un)substituted benzoylamino; R4 = H, OH, halo, CN, acyl, sulfamoyl, trialkylsioloxy, tetrasolyl, thienyl, pyrrolyl, pyrindinyl, oxazolyl, furanyl, or (un)substituted alkyl, alkenyl, alkymyl, alkoxy, amino, oxadiazolyl, Ph, pyridyl(oxy), carbamoyl; R11 and R12 = independently H, F, or C1 with the proviso that when one of R1 and R12 = F or C1, the other must be H; and pharmaceutically acceptable salts and esters thereof). The invention also relates to the use of I and their pharmaceutical compns. For treating hyperproliferative disorders and diseases associated with angiogenesis (no data). Examples include representative syntheses for compds. of the invention, pharmaceutical compns. comprising them, and tumor model assesys (no specific data given). For instance, N-Boc-indole was coupled with di-Me oxalate using t-Bull to give tert-Bu 2-[methoxy(oxolectyl]-1H-indole-1-carboxylate (72%). Cyclization of the dione with 1,2-phenylenediamine in AcOR afforded the quinoxalinone II (778). 694531-65-2P

PATENT NO. KIND DATE APPLICATION NO. DATE

US 2004087571 A1 20040506 US 2003-687015 20031016
W: AR, AG, AL, AM, AT, AU, AZ, BA, BB, BC, BR, BY, BZ, CA, CH, CN, CM, CO, CR, CU, CZ, DR, DK, DM, DZ, BC, ER, BY, BZ, CA, CH, CN, CM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LK, LS, LT, LU, LV, MA, NO, MM, KM, MM, MK, RM, NI, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SR, SB, ST, TJ, TM, TN, TR, TT, TZ, UA, UG, UG, UZ, VC, VN, YU, ZA, ZM, ZW
RM: GH, GM, KE, LS, MM, KZ, SD, SL, SZ, TZ, UG, ZM, ZM, AM, AZ, SY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, KE, SS, FI, FR, BG, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, TD, TG
AU 2003267800 A1 20040525 AU 2003-1B4614 W 20031020
R SOURCE(8): AU 2003267600 PRIORITY APPLN. INFO.: MARPAT 140:375191

The title compds. [I; R1 = (un)substituted heteroaryl; R2 = (un)substituted phenyl-(CH2)m-, naphthyl-(CH2)m-, cycloskyl-(CH2)m-, dkyl or heteroaryl-(CH2)m-; n = 0-4; R2 = H, (un)substituted alkyl,

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(Uass)
(antiproliferative and angiogenesis inhibitor; preparation of indolylquinoxalinones for treating hyperproliferative disorders and diseases essociated with angiogenesis)
694531-65-2 CAPLUS
6-Quinoxalinocarboxamide, 3-(3-amino-1H-indol-2-yl)-1,2-dihydro-N-(2-methoxyethyl)-N-methyl-2-oxo- (9CI) (CA INDEX NAME)

REFERENCE COUNT: THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 34 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCRSSION NUMBER: 2004:379708 CAPLUS
DOCUMENT NUMBER: 141:322446
TITLE: Picosecond time-resolved infrared investigation into

DOCUMENT NUMBER:

11.322446
Picosecond time-resolved infrared investigation into the nature of the lowest excited state of fac-[Re(cl)] (co) 3 (co2E-dppz)] (Co2E-dppz) (dipyrido[3,2a:2',3'c]) phenazine-11-carboxylic ethyl ester?

AUTHOR(S):

AUTHOR(S):

AUTHOR(S):

AUTHOR(S):

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CORPORATE SOURCE:

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Source:

Vibrational Spectroscopy (2004), 35(1-2), 219-223 (co2E-dpr)

COUNENT TYPS:

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AUTHOR photophysica of the fac [Re(Cl)(CO)3(CO2E-dppz)] (CO2E-dppz - dipyrido[3,2a:2',3'c]) phenazine-11-carboxylic Rt ester) was studied with picosecond time-resolved IR (ps-TRIR) spectras carbonyl (2100-1800 cm-1) and organic ester (1800-1600 cm-1) spectral regions. The ps-TRIR spectra in both regions give evidence for the formation of a metal-to-ligand charge transfer (MLCT) excited state. The magnitude of v(C.tplbond(O) shift in the metal carbonyl region of the excited state relative to chose of the ground state allow the excited state to be described more precisely as a d n(Re) + m (phenazine) 3MLCT

Tof330-96-99

RI: PEP (Physical, engineering or chemical process): PRP (Properties); PYP (Physical, process): SPN (Synthetic preparation): PRDS (Preparation): PRDC (Process)

(photophysics of tricarbonylchloro(dipyrido(phenazinecarboxylic Rt

cocess)
(photophysics of tricarbonylchloro(dipyrido[phenazinecarboxylic Rt
ester)rhenium complex studied with picosecond time-resolved IR

spectroscopy)
7-73-50-96-9 (ZDLUS
Rhenium, tricarbonylchloro(ethyl dipyrido[3,2-a:2',3'-c]phenazine-11-carboxylate-kN4,kN5)-, (OC-6-44)- (9CI) (CA INDEX NAME)

cycloalkyl-(CH2)n-, heterocycloalkyl-(CH2)n-, heteroaryl-(CH2)n-, aryl-(CH2)n-, n = 0-6; R3 and the carbon to which it is attached form (un) substituted and/or fused 5-7 membered carbocyclic ring; Y = heteroaryl, heterocycloalkyl, (un) substituted H3N-sulfonyl, C(:X)NH2; X = 0, S, (un) substituted H3N-sulfonyl, C(:X)NH2; X = 0, S, (un) substituted have substituted heterocycloalkyl-(CH2)p-, heteroaryl-(CH2)p-, phenyl-(CH2)p- or naphthyl-(CH2)p-; p = 0-4] which are CCR1 antagonists useful as immunomodulatory agents, were prepared E.g., a multi-step synthesis of quinoxaline-1-carboxylic acid [1(S)-benzyl-4(R)-benzylcarbasoyl-7-fluoro-2(S)-hydroxy-7-methyloctyllamide, was given. All of the compds. I that were tested showed ICS0 of <25 µM in the chemotaxis samey.

212789-34-39
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of heteroaryl-substituted heavanides.

(Uses)
(preparation of heteroary1-aubatituted hexanamides as CCR1 antagonists
useful as immunomodulatory agents)
21789-54-3 CAPLUS
6-Quinoxalinecarboxamide, N-{(18,28,48)-4-(aminocarbony1)-7-fluoro-2hydroxy-7-methy1-1-(phenylmethy1)octy1}- (9CI) (CA INDEX NAME)

LI3 ANSWER 36 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2003:991296 CAPLUS
TITLE: 140:41822
TRUENTOR(S): 140:41822
TRUENTOR(S): Allen, Darin; Fahr, Bruce; Oalob, Johan; Raimundo, Brian C.; Romanowski, Michael J.
SURGE: PATENT ASSIGNEE(S): Surgel Pharmaceuticals. Inc., USA
FOURCE: PCUBENT TYPE: Patent
LANGUAGE: PRIXED2
DOCUMENT TYPE: Patent
English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PA | TENT | NO. | | | KIN | D | DATE | | | APPL | ICAT | ION | NO. | | D. | ATE | |
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| NO | 2003 | 31035 | 99 | | A2 | | 2003 | 1218 | | WO 2 | 003- | US18 | 021 | | 2 | 0030 | 605 |
| MO | 2003 | 31035 | 99 | | A3 | | 2004 | 0708 | | | | | | | | | |
| | w: | AB, | AG, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BR, | BY, | BZ, | CA, | CH, | CN, |
| | | co, | CR, | CU, | CŹ, | DE, | DK, | DM, | DZ, | EC, | EE, | ES, | FI, | GB, | Œ, | GE, | GH, |
| | | GM, | HR, | Hυ, | ID, | IL, | IN, | 18, | JP, | KE, | KG, | KP, | ĸR, | KZ, | LC, | LK, | LR, |
| | | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NI, | NO, | ΝZ, | OM, |
| | | PH, | PL, | PT, | RO, | RU, | SC, | SD, | EE, | SG, | SK, | SL, | TJ, | TM, | TN, | TR, | TT, |
| | | TZ, | UA, | UG, | υz, | VC, | VN, | YU, | Zλ, | ZM, | ZW | | | | | | |
| | RN: | : GH, | GM, | KE, | LS, | MW, | MZ, | SD, | SL, | SZ, | TZ, | υα, | ZM, | ZW, | AM, | AZ, | BY, |
| | | KG, | ΚZ, | MD, | RU, | TJ, | TM, | AT, | BE. | BG, | CH, | CY, | CZ, | DE, | DK, | EE, | ES, |
| | | FI, | FR, | GΒ, | GR, | ΗU, | IE, | IT, | LU, | MC, | NL, | PT, | RO, | SE, | SI, | sĸ, | TR, |
| | | BP, | BJ, | CF, | CG, | CI, | CM, | GΑ, | GN, | GQ, | GW, | ML, | MR, | NE, | SN, | TD, | TG |
| ΑU | 2003 | 32389 | 48 | | A1 | | 2003 | 1222 | | AU 2 | 003- | 2389 | 8.8 | | 21 | 0030 | 605 |
| | NO
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N: AE,
CO,
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RW: GH,
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CO, CR,
GM, HR,
LS, LT,
PH, PL,
TZ, UA,
RM: GH, GM,
KG, KZ,
FI, FR, | MO 2003103599 MO 2003103599 M AE, AG, AG, CO, CR, CU, GM, HR, HU, LS, LT, LU, PH, PL, PT, TZ, UA, UG, RM: GH, GM, KE, KG, KZ, MD, FI, FR, GB, BF, BJ, CF, | MO 2003103599 A3 MC 2003103599 A3 CO CR. CU. CZ. GM. HR. HU. ID. LS. LT. LU. LV. PH. PL. PT. RO. TZ. UA. UO. KE. GH. GM. KE. LS. KO. KZ. MD. RJ. FF. FR. GB. GR. BF. BJ. CF. CO. | MO 2003103599 A2 MO 2003103599 A3 M: AE, AG, AL, AM, AT, CO, CR, CU, C2, DE, GM, HR, HU, ID, IL, LS, LT, LU, LY, MA, PH, PL, PT, RO, RU, TZ, UA, UG, UZ, VC, RM: GH, GM, KE, LS, MM, KO, KZ, MD, RU, TJ, FT, FR, GB, GR, HU, BF, BJ, CF, CO, CI. | MO 2003103599 A2 2003 MO 2003103599 A3 2004 MC AB, AG, AL, AM, AT, AU, CO, CR, CU, CZ, DB, DK, GM, HR, HU, ID, IL, IN, LS, LT, LU, LV, MA, MO, PH, PL, PT, RO, RU, SC, TZ, UA, UG, UZ, VC, VM, RM: GH, GM, KE, LS, MM, MZ, KO, KZ, MD, RU, TJ, TM, FT, FR, GB, GR, HU, IE, BP, BJ, CP, CQ, CI, CM, | MO 2003103599 A2 20031218 MO 2003103599 A3 20040700 M: AB, AG, AL, AM, AT, AL, AL, CO, CR, CU, CZ, DB, DK, DM, GM, HR, HU, ID, IL, IN, IS, LS, LT, LU, LV, MA, AD, MG, PH, PL, PT, RO, RU, SC, SD, TZ, LU, UG, UZ, VC, VN, TU, RM: GH, GM, KE, LS, MM, MZ, SD, KO, KZ, MD, RU, TJ, TM, AT, FT, FR, GB, GR, HU, IE, IT, BP, BJ, CF, CG, CI, CM, GA, | MO 2003103599 A2 20031218 MO 2003103599 A3 20040708 M: AE, AG, AL, AM, AT, AU, AZ, BA, CO, CR, CU, CZ, DE, DK, DM, DZ, GM, HR, HU, ID, IL, IN, IB, JP, LS, LT, LU, LV, MA, MD, MM, MC, PH, PL, PT, RO, RU, SC, SD, SE, TZ, LW, LG, LZ, VC, VN, VY, ZA, RM: GH, GM, KE, LS, MM, MZ, SD, SL, KG, KZ, MD, RU, TJ, TM, AT, BE, FI, FR, GB, GR, HU, IE, IT, LU, BP, BJ, CF, CG, CI, CM, GM, GM, | MO 2003103599 A2 20031218 MO 2 MO 2003103599 A3 20040708 M: AE, AG, AL, AM, AT, AU, AZ, BA, BB, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, GM, HR, HU, ID, IL, IN, IB, JP, KB, LS, LT, LU, LV, MA, MO, MO, MC, MN, PH, PL, PT, RO, RU, SC, SD, SE, SO, TZ, UA, UG, UZ, VC, VN, VY, UZ, AZ, AM, RM: GH, GM, KE, LS, MM, MZ, SD, SD, SZ, KG, KZ, MD, RU, TJ, TM, TB, BB, GG, FI, FR, GB, GR, HU, IE, IT, LU, MC, BP, BJ, CF, CG, CI, CM, AG, MI, AG, GB, | MO 2003103599 A2 20031218 MO 2003- MO 2003103599 A3 20040708 M: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, CO, CR, CU, CZ, DE, DK, DM, DZ, RC, EE, GM, RR, RU, ID, IL, IN, IS, JP, KE, KG, LS, IT, LU, LV, MA, MO, MO, KK, MN, MM, PH, PL, PT, RO, RU, SC, SD, ES, SO, SK, TZ, LUA, UG, UZ, UC, VN, YU, ZA, ZA, SA, RM1 GH, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, KG, KZ, MD, RU, TJ, TM, TB, BG, CC, FI, FR, GB, GR, HU, IE, IT, LU, MC, ML, BP, BJ, CP, CG, CI, CM, GA, GN, GO, GM, | MO 2003103599 A3 20031218 MO 2003-US18 M1 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, CO, CR, CU, CZ, DZ, OK, DM, DZ, KC, EK, ES, GM, RR, HU, ID, II, IN, IS, JP, KE, KG, KP, LS, LT, LU, LW, MA, MD, MG, MK, NR, MM, MG, PH, PL, TR, CR, US, CS, ED, ES, SG, SK, SL, TZ, LUA, UG, UZ, VC, VN, YU, ZA, ZM, ZM RM1 GH, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, KG, KZ, MD, RU, TJ, TM, AT, BR, BG, CH, CY, FI, FR, GB, GR, HU, IE, IT, LU, MC, NI, PT, SP, BJ, CF, CG, CI, CM, AG, AG, NG, OG, MF, ML. | MO 2003103599 A3 20031218 MO 2003-US18021 MO 2003103599 A3 20040708 M1 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CO, CR, CU, CZ, DE, OK, DM, DZ, KC, KE, ES, FI, GM, RR, HU, ID, II, IN, IS, JP, KE, KG, KP, KR, LS, LT, LU, LW, MA, AD, MG, MK, NR, MM, MK, MZ, PH, PL, PT, RG, RU, SC, SD, SE, SG, SK, SL, TJ, TZ, UA, UG, UZ, CV, VN, VU, ZA, ZM, ZM, KG, KZ, MD, RU, TJ, TM, AT, BS, BG, CM, CY, CZ, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RG, BP, BJ, CF, CG, CI, CM, AG, AG, NG, QG, MH, ML, MR, | MO 2003103599 A2 20031218 MO 2003-US18021 MO 2003103599 A3 20040708 M1: AB, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CO, CR, CU, CZ, DB, OK, DM, DZ, BC, EE, SS, FI, GB, GM, MR, HU, ID, II, NI, 18, JP, KE, KG, KP, KR, KZ, LS, LT, LU, LM, MA, MO, MO, MK, MN, MM, KK, MZ, NI, PH, PL, PT, RO, RU, SC, SD, EE, SG, BK, SL, TJ, TM, TZ, UA, UG, UZ, VC, VN, VU, ZA, ZM, ZM RM; GH, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, ZM, ZM, KG, KZ, MD, RU, TJ, TM, AT, BS, BG, CM, CY, CZ, DE, FI, FR, GB, GR, HU, IE, IT, LU, MC, NI, PT, RO, SR, SP, BJ, CF, CC, CI, CM, GA, GN, GQ, MH, LM, RN, SR, | MO 2003103599 A2 20031218 MO 2003-US18021 22 MO 2003103599 A3 20040708 M: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BC, BR, BY, BZ, CA, CO, CR, CU, CZ, DE, OK, DM, DZ, EC, EE, SS, FI, GB, GD, GM, HR, HU, ID, IL, IN, IS, JP, KS, KG, KP, KR, KZ, LC, LS, LT, LU, LW, MA, MC, MG, MK, MM, MK, KZ, AZ, MI, NO, PH, PL, PT, RO, RU, SC, SD, EE, BG, BK, EI, TJ, TM, TM, TZ, UA, UG, UZ, VC, VN, UZA, ZM, ZM RM; GH, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, ZM, ZM, KG, KZ, MD, RU, TJ, TM, AT, BS, BG, CM, CY, CZ, DS, DK, FI, FR, GB, GR, HU, IR, IT, LU, MC, NL, PT, RO, SR, ST, BF, BJ, CF, CC, CI, CM, GA, GN, GO, MH, LN, RN, SS, NS, | MO 2003103599 A2 20031218 MO 2003-US18021 20030 M1 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BC, BR, BY, BZ, CA, CH, CO, CR, CU, CZ, DE, OK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GM, HR, HU, ID, IL, IR, IB, JP, KS, KG, KP, KR, KZ, LC, LK, LS, LT, LUL, LY, MA, MC, MG, MC, MC, MM, MC, KZ, HI, MC, NZ, PH, PL, PT, RO, RU, SC, SD, EE, SG, SK, EE, TJ, TH, TR, TR, TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZM RM; GH, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, ZM, ZM, AA, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, ST, SK, SP, SJ, CF, CC, CI, CM, GA, QA, ON, GO, M, ML, NR, SS, SN, TD, |

US 2004048895 PRIORITY APPLN. INFO.:

A1 20040311

US 2003-456458 US 2002-386501P WO 2003-US18021

Compds. of formulas I and II [R1, R2 = aryl, sralkyl, heteroaryl, heteroarylkyl; L = linker] are prepared as caspase-1 inhibitors for the treatment of diseases such as inflammation, rheumatoid arthritis or sepsis. The compds. can also be used for preserving or storing mammalian organs or trisses by reducing apoptotic cell death. Thus, III was prepared in several steps.

S14203-31-1P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(Uses)
(preparation of acylamino(formyl)propanoic acids as caspase-1 inhibitors)
634203-91-1 CAPUS
Butanoic acid, 4-oxo-3-[[1-oxo-6-[(6-quinoxalinylcarbonyl)amino]-2-(2-thienyl)hexyl]amino]-, (38)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L13 ANSWER 37 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2003:836648 CAPLUS DOCUMENT NUMBER: 139:1350754 Preparation of 2,3-diphenvlouins

INVENTOR (S) :

119:350754
Preparation of 2,3-diphenylquinoxaline derivatives as inhibitors of Akt activity for treating cancer
Bilodeau, Mark T.; Duggan, Mark B.; Martnett, John C.; Lindsley, Craig W.; Manley, Peter J.; Wu, Zhicai; Zhao

PATENT ASSIGNEE(S):

PCT Int. Appl., 228 pp. CODEN: PIXXD2 Patent English 1 SOURCE :

DOCUMENT TYPE: LANGUAGE: PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

OTHER SOURCE(S): MARPAT 139:350754

The title compds. comprising a 2,3-diphenylquinoxaline moiety [I; u, v, w and x = CH, N; y, z = CH, N [provided that at least one of y and z = N); O = NNSSRS, (un) substituted aryl, heterocycly]; RI = alkenyl, halo, CN, etc.; R2 = OH, CN, CO2H, etc.; R3, R4 = H, alkyl, perfluoroalkyl; or R3 and R4 are combined to form (CH2)t wherein one of the carbon atoms is optionally replaced by O, SOm, (un) substituted NNCO, N(COH); RS, R6 = H, aryl, heterocyclyl, etc.; or NNSRs = monocyclic or bicyclic heterocycle; R7 = halo, CN, CO2H, etc.; n = 0-3; p = 0-2; t = 2-6; m = 0-2; q = 0-4; r = halo, CN, CO2H, etc.; n = 0-3; p = 0-2; t = 2-6; m = 0-2; q = 0-4; r = halo, CN, CO2H, etc.; n = 0-3; p = 0-2; t = 2-6; m = 0-2; q = 0-4; r = n-2; m = n-2;

(Uses)
(preparation of 2,3-diphenylquinoxaline derivs. as inhibitors of Akt
activity for treating cancer)
6:12847-31-3 CAPLUS
6-Ouinoxalinecarboxamide, 3-[4-[(4-(2,3-dihydro-2-oxo-1H-benzimidazol-1yl)-1-piperidiny|methyl|phenyl|-N-[3-(1H-imidazol-1-yl)propyl]-2-phenyl(9C1) (CA INDEX ANNEX)

612847-34-4 CAPLUS
6-Quinoxalinecarboxamide, 3-[4-[(4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yy)-1-piperidinyl]methyl]phenyl]-N-[3-(1H-imidazol-1-yl)propyl]-2-phenyl-bis(trifluoroacetate) (9C1) (CA INDEX NAME)

CM 1

CRN 612847-33-3 CMF C40 H38 N8 O2

PAGE 1-B

CRN 76-05-1 CMF C2 H F3 O2

612647-42-4 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-3-[4-[4-(2,3-dihydro-2-oxo-1H-benjinidazol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl- (9CI)
(CA INDEX NAME)

612847-43-5 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-2-[4-[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-3-phenyl- (9CI)
(CA INDEX NAME)

$$\mathtt{Et_2N-CH_2-CH_2-NH-C} \bigvee_{N}^{N} \bigvee_{Ph}^{CH_2-N} \bigvee_{N}^{N} \bigvee_{N}^{N}$$

612848-56-3 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-2-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl]-1-piperidinyl]methyl]phenyl]-3-phenyl-,
trifluoroacetate (9CI) (CA INDEX NAME)

CM 1

CRN 612847-43-5 CMF C40 H43 N7 O2

CM 2

CRN 76-05-1 CMF C2 H F3 O2

612848-57-4 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-3-[4-[4-(2,3-dihydro-2-oxo-1H-benzimidaxol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl-,
trifluoroacetate (9CI) (CA INDEX NAME)

CM 1

PAGE 1-A

612848-60-9 CAPLUS D-arabino-Hexose, 2-deoxy-2-[[[3-[4-[(4-(2,3-dihydro-2-oxo-1H-benzimidezol-1-y1)-1-piperidinyl]methyl]phenyl]-2-phenyl-6-quinoxalinyl]carbonyl]meino]-, (2ξ) - (9CI) (CA INDEX NAME)

CRN 76-05-1 CMF C2 H F3 O2

612848-58-5 CAPLUS
D-erabino-Rezose, 2-deoxy-2-{[[2-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazo1-1-y1)-1-piperidiny]]methyl]phenyl]-3-phenyl-6-quinoxalinyl]carbonyl]amino]-, (22)- (9C1) (CA INDEX NAME)

612848-59-6 CAPLUS
D-srabino-Hexoso, 2-deoxy-2-[[[2-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidszol-1-y1)-1-piperidinyl]methyl]phenyl]-3-phenyl-6-quinoxalinyl]carbonyl]amino]-, (25)-, trifluoroscetate (sait) [9C1] (CA INDEX NAME)

CRN 612848-58-5 CMF C40 H40 N6 O7

Absolute stereochemistry.

PAGE 1-B

612848-61-0 CAPLUS
D-arabino-Hexose, 2-deoxy-2-[[3-[4-[4-(2,3-dihydro-2-oxo-1H-benzimidezol-1-y])-1-piperidinyl]methyl]phenyl]-2-phenyl-6-quinoxalinyl]carbonyl]amino]-, (2\$)-, trifluoroacetate (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 612848-60-9 CMF C40 H40 N6 O7

CRN 76-05-1 CMF C2 H F3 O2

616868-43-0 CAPLUS 6-Quinoxalinecarboxamide, 2-[4-[[4-{2,3-dihydro-2-oxo-1H-benzimidazol-1-

yl)-1-piperidinyl]methyl]phenyl]-N-(3-(1H-imidazol-1-yl)propyl]-3-phenyl-(9CI) (CA INDEX NAME)

PAGE 1-B

RN 616866-44-1 CAPLUS
CN 6-Quinoxalinecarboxamide, 2-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-y1)-1-piperidinyl]methyl]phanyl]-N-[3-(1H-imidazol-1-y1)propyl]-3-phenyl-, trifluoroacetate (SCI) (CA INDEX RAME)

CM 1

CRN 616868-43-0 CMF C40 H38 N8 C2

PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 616868-46-3 CAPLUS
CN 6-Quinoxalinecarboxylic acid, 2-[4-[4-(2,3-dihydro-2-oxo-1H-bensimidazol-1-yl)-1-piperidinyllaethyl]phenyl]-3-phenyl-, ethyl ester, trifluoroacetate (9CI) (CA INDEX NAME)

CM 1

CRN 616868-45-2 CMF C36 H33 N5 O3

CV .

CRN 76-05-1

RN 616868-47-4 CAPLUS
CN 6-Quinoxalinecarboxylic acid, 3-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl-, ethyl ester (9CI) (CA INDEX NAME)

RN 616868-48-5 CAPLUS
CN 6-Quinoxalinecarboxylic acid, 3-{4-{[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-y]-1-piperidiny||methyl|phenyl|-2-phenyl-, ethyl ester, trifluoroacetate (9CI) (CA INDEX NAME)

СМ 1

CRN 616868-47-4 CMF C36 H33 NS O3

CH 2

CRN 76-05-1 CMF C2 H F3 O2

RN 616868-49-6 CAPLUS
CN 6-Quinoxalinecarboxamide, 2-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl}-N-(2-hydroxyethyl)-3-phenyl- (9Cl) (CA INDEX NAMS)

RN 616866-50-9 CAPLUS
CN 6-Quinoxalinecarboxamide, 2-[4-[]4-(2,3-dihydro-2-oxo-1H-benzimidezol-1-yl)-1-piperidinyl]methyl]phenyl]-N-(2-hydroxyethyl)-3-phenyl-trifluoroacetate (salt) [9C1] (CA INDEX RAME)

CH 1

CRN 616868-49-6 CMF C36 H34 N6 O3

$$\mathsf{HO-CH_2-CH_2-NH-C} \bigvee_{\mathsf{N}} \bigvee_{\mathsf{Ph}} \bigvee_{\mathsf{Ph}} \bigvee_{\mathsf{O}} \bigvee_{\mathsf{H}} \bigvee_{\mathsf{N}} \bigvee_{\mathsf{$$

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 616868-51-0 CAPLUS
CN 6-Quinoxalinecarboxamide, 3-[4-[[4-[2,3-dihydro-2-oxo-1H-benzimidazo]-1-yl]-1-piperidiny]]methyl]phenyl]-N-(2-hydroxyethyl)-2-phenyl- (9CI) (CA INDEX NAME)

RN 616868-52-1 CAPLUS
CN 6-Quinoxalinecarboxamide, 3-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1y1]-1-piperidinyl]methyl]phenyl]-N-(2-hydroxyethyl)-2-phenyl-,
trifluoroacetate (salt) (9CI) (CA INDEX NAME)

CM

CRN 616868-51-0 CMF C36 H34 N6 O3

CH 2

CRN 76-05-1 CMP C2 H F3 O2

616868-53-2 CAPLUS
6-Quinoxalinecarboxamide, 2-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperiddinyl]methyl]phenyl]-N-(2-hydroxy-2-phenylathyl)-3-phenyl-(SCI) (CA INDEX NAME)

616868-54-3 CAPLUS
6-Quinoxalinecarboxanide, 2-[4-[[4-{2,3-dihydro-2-oxo-1H-benzimidazol-1-yl]-1-piperidinyl]methyl]phenyl]-N-(2-hydroxy-2-phenylethyl)-3-phenyl-, trifluoroacetate (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 616868-53-2 CMP C42 H38 N6 O3

CM 2

CRN 76-05-1 CMF C2 H F3 O2

6-Quinoxalinecarboxamide, 3-[4-[{4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-y1)-1-piperidinyl]methyl]phenyl]-N-(2-hydroxy-2-phenylethyl)-2-phenyl-(9c1) (CA INDEX NAME)

616868-56-5 CAPLUS
6-Quinoxalinecarboxamide, 3-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]mathyl]phenyl]-H-(2-hydroxy-2-phenylethyl)-2-phenyl-, trifluoroacetate (salt) (9CI) (CA INDEX NAME)

CRN 616868-55-4 CMP C42 H38 N6 O3

CM 2

CRN 76-05-1 CMF C2 H F3 O2

616868-67-8 CAPLUS 6-Quinoxalinecarboxamide, 2-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidezol-1-yl)-1-piperidinyl]methyl]phenyl]-N-(5-hydroxypentyl)-3-phenyl- (9CI) (CA INDEX NAME)

616868-68-9 CAPLUS

6-Quinoxalinecarboxamide, 2-[4-[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-y1)-1-piperidinyl]methyl]phenyl-M-(5-hydroxypentyl)-3-phenyl-, trifluoroacetate (selt) (9CI) (CA INDEX NAME)

CM 1

CN 2

CRN 76-05-1 CMF C2 H F3 O2

616668-69-0 CAPLUS 6-Quinoxalinecarboxamide, 3-(4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-N-(5-hydroxypentyl)-2-phenyl- (9CI) (CA INDEX NAME)

616668-70-3 CAPLUS
6-Quinoxalinecarboxamide, 3-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidezol-1-yl)-1-piperidinyl]methyl]phenyl]-h-(5-hydroxypentyl)-2-phenyl-, trifluoroacetate (salt) (9CI) (CA INDEX RAMS)

CM 1

CRN 616868-69-0 CMF C39 H40 N6 O3

2

CRN 76-05-1 CMF C2 H F3 O2

RN 616868-71-4 CAPLUS
CN 6-Quinoxalinecarboxamide, 2-[4-[[4-{2,3-dihydro-2-oxo-1H-benzimidazol-1-y1}-1-piperidinyl]methyl]phenyl]-N-[3-(dimethylamino)-2,2-dimethylpropyl]-3-phenyl- (9CI) (CA INDEX NAME)

PAGE 1-B

616868-72-5 CAPLUS
6-Quinoxalinecarboxamide, 2-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1y]]-1-pjerdidnyl]methyl]phenyl]-N-[3-(dimethylamino)-2,2-dimethylpropyl]3-phenyl-, trifluoroacetate (9CI) (CA INDEX NAME)

CM 1

CRN 616868-71-4 CMF C41 H45 N7 O2

PAGE 1-B

CRN 76-05-1 CMF C2 H F3 O2

616868-73-6 CAPLUS
6-Quinoxalinecarboxamide, 3-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1yl)-1-piperidinyl]methyl]phenyl]-N-[3-(dimethylamino)-2,2-dimethylpropyl]2-phenyl- (9CI) (CA INDEX NAME)

PAGE 1-A

RN 616868-74-7 CAPLUS

PAGE 1-B

616868-76-9 CAPLUS
6-Quinoxalinecarboxamide, 2-[4-([4-(2.3-dihydro-2-oxo-1H-benzimidezol-1yl)-1-piperidinyl]methyl]phenyl]-N-(3-hydroxy-2,2-dimethylpropyl)-3-phenyl-, trifluoroacetate (salt) [9CI] (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

CRN 76-05-1 CMP C2 H F3 O2

616866-77-0 CAPLUS
6-Quinoxalinecarboxamide, 3-{4-[{4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-N-(3-hydroxy-2,2-dimethylpropyl)-2-phenyl-(9CI) (CA INDEX NAME)

6-Quinoxalinecarboxamide, 3-[4-[[4-{2,3-dihydro-2-oxo-1H-benzimidazol-1-yl]-1-piperidinyl]methyl]phenyl]-N-[3-(dimethylamino)-2,2-dimethylpropyl]-2-phenyl-, trifluoracetate (8C1) (CA INDEX RAME)

CM 1

CRN 616868-73-6 CMF C41 H45 N7 O2

PAGE 1-B

CRN 76-05-1 CMF C2 H F3 O2

616868-75-8 CAPLUS
6-Quinoxalinecarboxamide, 2-[4-[[4-(2,3-dihydro-2-oxo-1H-benrimidazol-1yl)-1-piperidinyl]methyl]phenyl]-N-(3-hydroxy-2,2-dimethylpropyl)-3-phenyl(9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

616868-78-1 CAPLUS
6-Quinoxalinecarboxamide, 3-[4-[[4-[2,3-dihydro-2-oxo-1H-benzimidazol-1-y]]-1-pjeridinyl]methyl]phenyl]-N-[3-hydroxy-2,2-dimethylpropyl]-2-phenyl-trifluoroacetate (malt) (9CI) (CA INDEX NAME)

CRN 616868-77-0 CMF C39 H40 N6 O3

PAGE 1-B

CH 2

CRN 76-05-1 CMF C2 H F3 O2

616868-79-2 CAPLUS
6-Quinoxalinecarboxamide, N-[[4-(aminosulfonyl)phenyl]methyl]-2-[4-[[4-(2,3-dhydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl)phenyl]-3-phenyl-[9CI) (CA INDEX NAME)

PAGE 1-B

616868-80-5 CAPLUS
6-Quinoxalinecarboxamide, N-[[4-(aminosulfonyl)phenyl]methyl]-2-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-3-phenyl-, trifluoroacetate (9CI) (CA INDEX NAME)

CM 1

CRN 616868-79-2 CMF C41 H37 N7 O4 8

CRN 76-05-1 CMF C2 H F3 O2

616868-81-6 CAPLUS
6-Quinoxalinecarboxamide, N-[[4-(aminosulfonyl)phenyl]mathyl}-3-[4-[[4-(2,3-dihydro-2-oxo-1H-benzinidazol-1-yl)-1-piperidinyl]mathyl]phenyl]-2-phenyl-(SCI) (CA INDEX NAME)

PAGE 1-B

616868-82-7 CAPLUS
6-Quinoxalinecarboxamide, N-[[4-(aminosulfonyl)phenyl]methyl]-3-[4-[[4-(2,3-dh)ydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl-, trifluoroacetate (9CI) (CA INDEX NAME)

CM 1

CRN 616868-81-6 CMF C41 H37 N7 O4 S

PAGE 1-B

CRN 76-05-1 CMF C2 H F3 O2

616869-39-7 CAPLUS
6-Quinoxalinecarboxamide, 3-(4-[[4-(1H-benzimidezol-2-yl)-1piperidinyl]methyl]phenyl]-N-(2-hydroxyethyl)-7-methyl-2-phenyl(CA INDEX NAME)

616669-40-0 CAPLUS
6-Quinoxalinecarboxamide, 3-[4-[[4-(1H-benximidazol-2-yl]-1-piperidinyl]methyl]phenyl]-N-(2-hydroxyethyl)-7-methyl-2-phenyl-, trifluoroacetate (salt) (9CI) (CA INDEX NAME)

CRN 616869-39-7 CMF C37 H36 N6 O2

2

CRN 76-05-1 CMF C2 H F3 O2

RN 616869-41-1 CAPLUS
CN 6-Quinoxalinecarboxamide, 2-[4-[(4-(1H-benzimidazol-2-yl)-1-piperidinyl)methyl]phenyl]-N-(2-hydroxyethyl)-7-methyl-3-phenyl-(CA INDEX NAME)

RN 616869-42-2 CAPLUS
CN 6-Quinoxalinecarboxamide, 2-{4--[{4--(14--benzimidazol-2-y1)-1-piperidinyl]=sethyl]phenyl]-N-(2--hydroxyethyl)-7-methyl-3-phenyl-,
trifluoroacetate (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 616869-41-1 CMF C37 H36 N6 O2

CM 2

CRN 76-05-1

RN 616869-89-7 CAPLUS
CN 6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-3-[4-[4-(2-methyl-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

RN 616869-90-0 CAPLUS
CN 6-Quinoxalinecarboxamide, N-(2-hydroxyethyl)-3-[4-[(4-(2-methyl-1H-benximidazol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl- (9CI) (CA INDEX NAME)

RN 616869-92-2 CAPLUS
CN 6-Quinoxalinecarboxamide, N-{3-{1H-imidazol-1-yl}propyl}-3-{4-{{4-(2-methyl-1H-benzimidazol-1-yl}-1-piperidinyl]methyl]phenyl}-2-phenyl- (9CI)
(CA INDEX NAMS)

PAGE 1-

RN 616869-95-5 CAPLUS
CN 6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-3-{4-[4-(5-fluoro-1H-benzimidazol-2-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl- (9CI) (CA INDEX NAUE)

PAGE 1-A

PAGE 1-B

RN 616870-60-1 CAPLUS
CN 6-Quinoxelinecarboxamide, 3-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yp)-piperidinyl]methyl]phenyl]-N-{3-(1H-imidazol-1-yl)propyl]-2-phenyl-, trifluoroacetate (9CI) (CA INDEX NAME)

CM 1

CRN 612847-33-3 CMP C40 H38 N8 O2

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PAGE 1-B

CM :

CRN 76-05-1 CMF C2 H F3 O2

RN 616870-81-6 CAPLUS
CN 6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-1-[4-[[4-(2-methyl-1H-bensimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl-,
trifluoroacetate (9CI) (CA INDEX NAME)

CM 1

CRN 616869-89-7 CMF C41 H45 N7 O

PAGE 1-B

CM 2

CRN 76-05-1 CMP C2 H F3 02 E-C-CO3H

RN 616870-82-7 CAPLUS
CN 6-Quinoxalinecarboxamide, N-(2-hydroxyethyl)-3-[4-[4-(2-methyl-lH-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl-, trifluoroacetate (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 616869-90-0 CMF C37 H36 N6 O2

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 616870-84-9 CAPLUS
CN 6-Quinoxalinecarboxamide, N-[3-(1H-imidezol-1-yl)propyl]-3-[4-{[4-(2-methyl-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl-, trifluoroacetate (9CI) (CA INDEX NAME)

CH 1

CRN 616869-92-2 CMF C41 H40 N8 O

CRN 76-05-1 CMF C2 H F3 O2

F-C-002H

616870-87-2 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-3-[4-[[4-(5-fluoro-1H-benzinidzol-2-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl-, trifluoroacetate (9CI) (CA INDEX NAME)

CRN 616869-95-5 CMF C40 H42 F N7 O

PAGE 1-A

CRN 76-05-1 CMF C2 H F3 O2

homol. domain of Akt. Thus, 3,6-dichloropyridazine was converted to its
4-cyclobutyl derivative which was cyclized with BENNINN2 and aminated to give I
[R1 = Ph, R2 = NHCH2CMe2CH2NMe2, R3 = H, R4 = cyclobutyl]. This compound
had ICSO for inhibition of Akt 1 of 1.4 pm.
612847-34-49 612847-42-49 612847-43-59
612848-66-19 612248-37-49 612848-59-69
612848-61-09
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
(preparation of triazolo(4,3-b) pyridazines and 2,3-diarylquinazolines for
the treatment of cancer)
612847-34-4 CAPLUS
6-Quinoxalinecarboxamide, 3-[4-[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1yl)-1-piperidinyl]methyl]phenyl]-N-(3-(1H-imidazol-1-yl)propyl]-2-phenyl-,
bis(trifluoroacetate) (9CI) (CA INDEX NAME)

CM 1

CRN 612847-33-3 CMF C40 H38 N8 O2

PAGE 1-A

г-с-со₂н

612847-42-4 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-3-[4-[[4-(2,3-dihydro-2-oxo-1H-benrimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl- (9CI)

REFERENCE COUNT:

L13 ANSWER 36 OF 161 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC, NUM. COUNT:

| PATENT INFOR | MATT | UN: | | | | | | | | | | | | | | |
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| WO 2003 | 0844 | 73 | | A2 | | 2003 | 1016 | | WO 2 | 003- | US10 | 632 | | 2 | 0030 | 404 |
| NO 2003 | 0844 | 73 | | A3 | | 2004 | 0212 | | | | | | | | | |
| W: | AE, | AG, | AL, | AM, | AT, | AU, | AZ, | BA, | 88, | BG, | BR, | BY, | BZ, | CA, | CH, | CN, |
| | ço, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC, | EE, | ES, | FI, | GB, | GD, | GE, | GH, |
| | GM, | HR, | Hυ, | ID. | IL, | IN, | IS, | J₽, | KE, | KG, | KR, | KZ, | LC, | LK, | LR, | LS, |
| | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NI, | NO. | NZ, | OM, | PH, |
| | PL, | PT. | RO, | RU, | SC. | SD. | SE. | SG, | SK. | SL. | TJ. | TM. | TN, | TR, | TT, | TZ. |
| | UA, | UG. | US. | UZ. | VC. | VN, | YU. | ZA, | ZM. | ZW | | | | | | |
| RW: | GH, | GM, | KE, | LS, | MN, | MZ, | SD, | SL, | SZ, | TZ, | UG, | ZM, | ZW, | AM, | AZ, | BY, |
| | KG. | KZ, | MD, | RU, | TJ. | TM, | AT, | BE, | BG, | CH, | CY, | CZ, | DE, | DK. | EE, | ES, |
| | PI. | FR. | GB, | GR. | HU, | IE, | IT. | LU. | MC. | NL. | PT. | RO, | SE, | SI. | SK, | TR. |
| | BF, | ВJ, | CF, | CG, | CI, | CM, | GA, | GN, | GQ, | GW, | ML, | MR, | NE, | SN, | TD, | TG |
| AU 2003 | 2263 | 01 | | A1 | | 2003 | 1020 | - 1 | AU 2 | 003- | 2263 | 01 | | 2 | 0030 | 404 |
| PRIORITY APP | LN. | INFO | . : | | | | | 1 | US 2 | 002- | 3708 | 27P | | P 2 | 0020 | 408 |
| | | | | | | | | | US 2 | 002- | 4172 | 02P | 1 | P 2 | 0021 | 009 |
| | | | | | | | | 1 | NO 2 | 003-1 | US10 | 632 | 1 | W 2 | 0030 | 404 |
| ~* | | | | | | | | | | | | | | | | |

GI

Triezolo[4,1-b]pyridazines I [R1 = (un)substituted Ph, furyl, thienyl, pyridinyl; R2 = substituted NH2, OH; R3 = H, R4 = (un)substituted cycloslkyl, aryl; R8R4 = (un)substituted GH:GHCH:GH) and quinazolines II [R5, R6 = (un)substituted Ph; R7 = H, alkyl, halogen, GH, alkoxyl were prepared for use as sinhibitors of one or two of the isoforms of Akt, a serine/threonine protein kinase, acting particularly on the pleckstrin

612847-43-5 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-2-{4-[4-(2,3-dihydro-2-oxo-1R-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl)-3-phenyl- (9CI)
(CA INDEX NAME)

612848-56-3 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-2-[4-[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-3-phenyl-, trifluoroacetate (9CI) (CA INDEX NAME) RN CN

CRN 612847-43-5 CMF C40 H43 N7 O2

2

CRN 76-05-1 CMF C2 H F3 O2

612648-57-4 CAPLUS
6-Quinoxalinecarboxamide, N-[2-(diethylamino)ethyl]-3-[4-[4-(2,3-dihydro-2-oxo-1H-benzinidazol-1-yl)-1-piperidinyl]methyl]phenyl]-2-phenyl-,
trifluoroacetate (9CI) (CA INDEX NAME)

CRN 612847-42-4 CMF C40 H43 N7 O2

CH 2

RN 612848-59-6 CAPLUS
CN D-arabino-Hexose, 2-deoxy-2-[[[2-[4-[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-yl)-1-piperidinyl]methyl]phenyl]-3-phenyl-6-quinoxalinyl]carbonyl]amino]-, [22]-, trifluoroacetate (salt) [901] (CA INDEX NAME)

CM 1

Absolute stereochemistry.

L13 ANSWER 39 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2003:677647 CAPLUS
DOCUMENT NUMBER: 1003:677647 CAPLUS
TITLS: 149319 enhanced duplex etability of dipyrido(3,2-a:2',3'-c)phenazine-modified oligonuclectide conjugate
AUTHOR(S): Kitamura, Yusuke; Chara, Toehihiro; Shirasaka, Yoshinori; Mitsuru, Tomonori; Tazaki, Masato; Jyo, Akinori
CORPORATE SOURCE: Department of Applied Chemistry and Biochemistry, Kumanoto University, Kumanoto, 860-8555, Japan Nucleic Acide Research Supplement (2003), 3(3rd International Symposium on Nucleic Acide Chemistry (and) 30th Symposium on Nucleic Acide Chemistry in Japan, 2003), 95-96
CODEN: NARSCE
OCCURNY TYPE: Journal Acide Chemistry Press
DOCUMENT TYPE: Journal English
AB Dipyrido(3,2-a:2',3'-c)phenazine (DPPZ) or 1,10-phenanthroline (Phen) was tethered to the 5'-end of a short oligonucleotide (ODN) to generate two ODN conjugates. The conjugates formed stable duplexes with complementary 6 mer (d(TTAGOG)), which is one unit of telomeric repeate of human. The of the corresponding duplex with unmodified 6 mer by 19.6 °C. This stabilization is enormous compared with those observed in other ODN conjugates reported previously. It would be attributed to the effective interaction of tethered heterosrom, groups with DNA base stack of the duplex-19-8P
RL: PRP (Propertice): SPN (Symthetic preparation): PREP (Preparation) (Dreparation and thermodon, of highly enhanced duplex stability of dipyridophenazine-modified oligodeoxyribonucleotide conjugates)
RN 563942-79-8 CAPLUS
CN Guanosine, thymidylyl-(3'-5')-2'-deoxyquanylyl-(3'-5')-2'-deoxyquanylyl-(3'-5')-2'-deoxyquanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'-deoxyydanylyl-(3'-5')-2'

CM 1

CRN 663942-78-7 CMP C82 H96 N26 O37 P6

Absolute stereochemistry.

612848-61-0 CAPLUS
D-arabino-Hexose, 2-deoxy-2-[[[3-[4-[[4-(2,3-dihydro-2-oxo-1H-benzimidazol-1-y]]-1-piperidinyl]methyl]phenyl]-2-phenyl-6-quinoxalinyl]carbonyl]amino]-, (2\$)-, trifluoroacetate (selt) (9CI) (CA INDEX NAME)

CRN 612848-60-9 CMF C40 H40 N6 O7

PAGE 1-B

PAGE 1-B

CRN 117490-04-7 CMF C60 H75 N24 O35 P5

Absolute stereochemistry.

PAGE 1-A

OTHER SOURCE(S):

WO 2003-SE258 CASREACT 139:197375; MARPAT 139:197375

CR2R3 (CH2) mCR4 (OH) CR5R6 (CR7R8) nNR32 ZYR

The invention provides piperidinyl alcs. (shown as I; variables defined below; e.g. N-[(2R)-1-(4-(3,4-dichlorophenoxy)piperidin-1-y1]-2-hydroxypropyl)-2-(methylaulfonyl)benzamide) for use as modulators of chemokine receptor (sepecially CR3) activity for use in, for example, treating asthma. For I: X is CH2, 0, 8(0)2 or NRIO; Y is a bond, CH3, NR35, CH3NN, CH3NNC(O, CH(OH), CH(NROCA3)), OK (NRHOGA7)4, CH3O or CH28; Z is C(0), or when Y is a bond Z can also be S(0)2; R1 is (un)substituted aryl, (un)substituted heterocycly) or C4-6 cyclocally [used to a benzene ring; addnl. details are given in the claims. Percent inhibition at 3 nM cotaxin of cotaxin-nediated human cosinophil chemotaxis is tabulated for 16 examples of 1, e.g. 106 % for N-[(2R)-3-[4-(3,4-dichlorophenoxy)piperidin-1-y1]-2-hydroxypropyl]-1-oxo-1,2-dinydroisoquinoline-4-carboxamide. Histamine H1 receptor binding activity was datermined for the same compdex., e.g. pKi = 8.4 for N-[(2R)-3-[4-(3,4-dichlorophenoxy)piperidin-1-y1]-2-hydroxypropyl]-1-oxo-1,2-dinydroisoquinoline-4-carboxamide. 48 Example prepns. of intermediates and 334 of I are included. For example, to propare N-[(2R)-3-[4-(3,4-Dichlorophenoxy)piperidin-1-y1]-2-hydroxypropyl]-3-(mathylaulfonyl)benzamide (0.055 g), a mixture of 2-(mathylaulfonyl)benzolc

PAGE 2-B

REFERENCE COUNT:

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 40 OF 181 CAPLUS ACCESSION NUMBER: 2003: DOCUMENT NUMBER: 139:17

APLUS COPYRIGHT 2006 ACS on STN
2003:656742 CAPLUS
139:197375
Preparation of piperidinyl alcohols as chemokine
receptor modulators for treatment of diseases such as
asthma
Alcaras, Lilian; Purber, Mark; Purdie, Mark;
Springthorpe, Brian
Astrazeneca A.B., Swed.
PCT Int. Appl., 166 pp.
CODEN: PIXKD2
Patent INVENTOR (S) .

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE:

acid (0.063 g), (2R)-1-amino-3-[4-(3,4-dichlorophenoxy)piperidin-1-yl]propan-2-ol (0.1 g) and N,N-diisopropylethylamine (0.1 mL) in dry DMF (3 mL) was cooled to 0° with stirring; 2-(1H-9-azabenzotrizato-1-yl]-1,1,3,-1-teramethyluronium hexafluorophenoxylosphate (0.13 g) was added and the mixture was stirred at 0° for 1-2 h. The invention also provides a process for making 4-(3,4-dichlorophenoxyl)piperidine, which is useful as an intermediate for making certain compds. of the invention. The process comprises (a) reacting 4-hydroxylperidine with a suitable base in a suitable solvent at room temperature; and (b) heating the mixture so produced

1,2-dichloro-4-fluorobenzene at 50-90°, or at reflux of the solvent

used.
\$583892-05-7P, N-[(2R)-3-[4-(3,4-Dichlorophenoxy)piperidin-1-yl]-2hydroxypropyl]quinoxaline-5-carboxamide
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USSS

(Therapeutic use; size (transparent).
(Uses)
(drug candidate; preparation of piperidinyl alcs, as chemokine receptor modulators for treatment of diseases such as asthma)
583882-05-7 CAPLUS
6-Ouinoxalinecarboxamide, N-[(2R)-3-[4-(3,4-dichlorophenoxy)-1-piperidinyl]-2-hydroxypropyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

INVENTOR (S):

L13 ANSWER 41 OF 181 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

THERE ARE 11 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 2001:376619 CAPLUS COPYRIGHT 2006 ACS on STN 2001:376619 CAPLUS Preparation of N.N'-substituted-1,3-diamino-2-hydroxypropense for treating Althouser's disease Varghese, John: Maillard, Michel; Jagodzinske, Berbara; Beck, James P.; Gallunes, Andrea; Fang, Lerry; Seely, Jennifer; Tenbrink, Ruth; Freskos, John; Mickelson, John; Semale, Lakshman; Mom. Roy Elen Pharmaccuticals, Inc., USA; Pharmacia 6 Upjohn Company PCT Int. Appl., 1243 pp. CODEN: PIXXD2 Patent English 2

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT ASSIGNEE(S):

| PAT | ENT | NO. | | | KIN | D | DATE | | | appl | ICAT | ION . | NO. | | D. | ATE | |
|-----|------|------|-----|-----|-----|-----|------|------|-----|------|------|-------|-----|-----|-----|------|-------|
| | | | | | | • | | | | | | | | | - | | • • • |
| WO | 2003 | 0400 | 96 | | A2 | | 2003 | 0515 | | NO 2 | 002- | U836 | 072 | | 21 | 0021 | 108 |
| WO | 2003 | 0400 | 96 | | A3 | | 2004 | 0506 | | | | | | | | | |
| | W: | AE, | AG. | AL, | AM, | AT, | AU. | AZ, | BA, | BB, | BG, | BR, | BY, | BZ, | CA, | CH, | CN |
| | | co. | CR. | CU. | CZ. | DE. | DK. | DM. | DZ, | EC, | EE, | ES, | FI, | GB, | GD, | GE, | GH |
| | | GM. | HR. | HU. | ID. | IL. | IN. | IS, | JP. | KE, | KG, | KP, | KR, | KZ, | LC, | LK, | LR |
| | | LS. | LT. | LU. | LV. | MA. | MD. | MG, | MK. | MN. | MW. | MX, | MZ, | NO. | NZ, | OM, | PH |
| | | PL. | PT. | RO. | RU. | SD. | SE. | SG. | SI. | SK. | SL. | TJ, | TM, | TN, | TR, | TT, | TZ |
| | | | | | | | YU, | | | | | | | | | | |

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2, SK
20021108
20021108
20021108
20040511
20040507
P 20011108
P 20011228
P 20020103
W 20021108
                                                                                  AL, TR, BG, CZ, BR 2002-14035
JP 2003-542142
CN 2002-826786
ZA 2004-3578
NO 2004-3359
US 2001-344086P
US 2002-345635P
JP 2005520791
CN 1759095
ZA 2004003578
NO 2004002359
PRIORITY APPLN. INFO.:
                                                                                    WO 2002-US36072
OTHER SOURCE(S):
                                               MARPAT 138:385173
```

The title compds. [I; R1 = (un)substituted alkyl, alkenyl, alkynyl, etc.; R2 = H, alkyl, haloalkyl, alkenyl, etc.; R3 = H, alkyl, haloalkyl, alkenyl, etc.; R3 = H, alkyl, haloalkyl, alkenyl, etc.; or R2 and R3 are taken together with the carbon to which they are attached to form a carbocycle of 3-7 carbon atoms, optionally where one carbon atom is replaced by a heterostom selected from the group consisting of O, 8, SO2, (un)substituted NH; R4 = alkyl, haloalkyl,

UA. UG. US. UZ. VC. VN. YU. ZA. ZM. ZM
RM: GH. GM. KE, LE, MM. MZ. SD, SL. SZ. TZ. UG, ZM. ZM. AM. AZ. BY,
KG. KZ. MD. RU, TJ. TM. AT. BE. BG, CH. CY. CZ. DE, DK. EE, ES,
FI. FR. GB, GR, IE, IT. LU, NC. NL, FT, SE, SK. TR. BF. BJ. CF.
CG. CI. CM. GA, GN, GO, GM, ML, MR, NE, NS, NT, D. TG
EP 1439169
R: AT. BE, CH. DE, DK. ES, FR, GB, GR, IT. LI, U, NL, SE MC. PT.
IE, SI, LT. LV, FI, RO, MK, CY, AL. TR, BG, CZ, EE, SK
US 2004248872
A1 20041209
PRIORITY APPLN. INFO::

US 2002-2995687
JP 2002-1-2995687
MARPAT 138:304304

COTHER SOURCE(S):

MARPAT 138:304304

Dr 2001-299587 A 20010928

JP 2002-142199 A 20010921

JP 2002-142199 A 20020930

OTHER SOURCE(S):

MARPAT 138:304304

AB The difluoroalkenyl heterocyclecarboxylate, -thiocarboxylates, or dithiocarboxylate derive. represented by the general formula O-C(:L1)-L2-(CH2)n-C(CF3):CP2 or pharmacol. acceptable salts thereof (wherein Li and L2 are the same or different and each represents oxygen or sulfur; n is an integer of 2 to 8; and O represents an optionally substituted 5 - to 12-membered heterocyclic group having any desired heterostom selected among nitrogen, oxygen, and sulfur wherein the heterostom in the heterocyclic ring is a nitrogen, it may be oxidized to N-oxide), which are useful as insecticides, carricides, and nematocides, are prepared These compda. are sufficiently effective in controlling various pests even when used in a small dose and are highly safe for crops, natural enesies to the pests, and animals. Thus,
4-phenyl-1,2,3-thiadiazole-5-carboxylic acid 0.23, 6,6-difluoro-5-methyl-5-hexenol 0.17, and 4-dimethylaminopyridine 0.13 g were dissolved in 4 mj.
CH2C12, treated with 0.29 g 1-ethyl-3-(13-dimethylaminoproyl)carbodifinide hydrochloride at room temperature, and stirred for 20 h to give 6,6-difluoro-5-methyl-5-hexenyl 4-phenyl-1,2,3-thiadiazole-5-carboxylate (I). I and 4,4-difluoro-1-methyl-3-bethyl-1-6-thenyl-6-betoxyl-2-methylpyrimidine-4-carboxylate at 500 pps controlled 2904 4th instar larvae of Nilaparvata lugens.

IT 509100-31-69
R: ARR (Agricultural use); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of difluoroalkenyl heterocyclecarboxylate and -thiocarboxylate as pest control asents and an animals.

(Uses)
(preparation of difluoroalkenyl heterocyclecarboxylate and -thiocarboxylates as pest control agents such as insecticides, acaricides, and nematocides)
509100-31-6 CAPLUS
6-Ouinoxalinecarboxylic acid, 4,4-difluoro-3-methyl-3-butenyl sster (9CI)
(CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 43 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2002:889219 CAPLUS
DOCUMENT NUMBER: 137:379407
Colorimetric sensor compositions and methods
INVENTOR(S): Sessler, Jonathan; Andriolstti, Bruno; Try, Andrew
Carl; Black, Christopher
DATENT ASSIGNEE(S): USA

U.S., 30 pp. CODEN: USXXAM

hydroxyalkyl, etc.; RS = R6X (wherein X = CO, SO2, (un)substituted CH2; R6 = (un)substituted Ph, naphthyl, indanyl, etc.); R15 = H, alkyl, alkoxy, etc.) which have activity as inhibitors of B-secretaes and ars therefore useful in treating a variety of disorders such as Altheimer's disease, were prepared E.g., a multi-step synthesis of (18,7R)-11, starting from (28)-2-[(tert-butoxycarbonyl)aminol-1-(1,5-difluorophenyl)propanoic acid, was given. The compds. I showed ICSO of < 20 µM in cell free inhibition assay utilizing a synthetic APP substrate. This is a Part 1 of

Inhibition assay utilizing a synthetic APP substrate. This is a vart 10.
1-2 series.
327723-62-8P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); TRU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of N.N'-substituted-1,3-diamino-2-hydroxypropanes for treating Alzheiner's disease)
527735-62-8 CAPUS
6-Quinoxalinecarboxamide, N-[(18,2R)-1-[(3,5-difluorophenyl)methyl]-3-[(3-ethylphenyl)methyl]amino]-2-hydroxypropyl]-1,2,3,4-tetrahydro-2,3-dioxo-(9CI) (CA INDEX NAME)

L13 ANSWER 42 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER: 2001:282533 CAPLUS
DOCUMENT NUMBER: 138:304304
TITLS: Preparation of 445

130:304304 Preparation of difluoroalkene derivatives as pest control agents containing the same, and intermediate therefor

therefor
Abo, Tetsuya; Tamai, Ryuji; Ito, Minoru; Tamaru,
Mseatoshi; Yano, Hiroyuki; Tekahashi, Satoru;
Muramatsu, Norimichi
Kumiai Chemical Industry Co., Ltd., Japan; Ihara
Chemical Industry Co., Ltd.
PCT Int. Appl., 195 pp.
CODEN: PIXXD2
Patent INVENTOR (S):

PATENT ASSIGNEE(S):

SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

ENT NO. KIND DATE APPLICATION NO. DATE

2003029211 A1 20030410 NO 2002-JP10142 20020930
N: AS, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GB, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MG, MG, MK, MN, MN, MX, MZ, MO, NZ, MM, PL, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TH, TN, TR, TT, TZ, PATENT NO. WD 2003029211

DOCUMENT TYPE: LANGUAGE: PAMILY ACC. NUM. COUNT: PATENT INFORMATION: Patent English

B1 20021119 A1 20030828 PATENT NO. APPLICATION NO. DATE US 2000-579040 US 2002-222028 US 1999-136467P US 2000-579040 US 6482949 US 2003162960 PRIORITY APPLN. 1NFO.:

WS 1797-134657P P 19990528 US 2000-579040 A3 20000526

R SOURCE(S): MARPAT 137:379407 The present invention provides novel compds. exemplified by pyrrolic nitrogens used as anion and neutral species recognition elements with an aromatic core as a signal group. Described are methods for the synthesis of various pyrrole aryl compds. as well as various applications for these compds. Methods of use include the binding and detection of specific analytes in a mixture and, in some examples, the separation of the analyte from the mixture Addml. methods of use include the transport of therapeutic agents and the sensing of components, degradants, and impurities in foodstuffs. OTHER SOURCE(S): ΙT

foodstwffs.
475476-81-4P
RL: SPN (Synthetic preparation); PREP (Preparation)
(colorimetric sensor compss. and methods based on pyrrole-aryl compds.
for anion and neutrous recognition and determination)
475476-81-4 CAPUS
6-Quinoxalinecarboxylic acid, 2,3-di-1H-pyrrol-2-yl-, octyl ester (9CI)
(CA INDEX NAME)

REFERENCE COUNT: THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 44 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1002:808500 CAPLUS
115:34844

AUTHOR(S): SOURCE: 118:14844

AUTHOR(S): CORPORATE SOURCE: 118:14849

SOURCE: SOURCE: 124(45), 13416-13433

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

CODEN: JACSAT; ISSN: 0002-7863 American Chemical Society Journal

CODEN: JACSAT: ISSN: 0002-7863

American Chemical Society

Journal
LANGUAGE:

DOURNANT TYPE:

JOURNAI

AB Me here report our studies on the conjugation of photoreactive Ru2+

complex to oligonucleotides (ODNs), which give a stable duplex with the

complementary target DNA strand. These functionalised DNA duplexes

bearing photoreactive Ru2- complex can be specifically Dhocolyyard to give

the strand of th

CRN 16919-18-9 CMF P6 P CCI CCB

478819-70-4 CAPLUS
Ruthenium(2+), [N-[2-(2-(2-3-dihydroxypropoxy)ethoxy]

CH 1

CRN 478819-69-1 CMP C48 H45 N9 O6 Ru CCI CCS

PAGE 1-A

478818-94-9 CAPLUS Ruthenium(2+), (acetonitrile) (N-ethyldipyrido[3,2-a:2',3'-c]phenazine-l1-carboxanide-KM, kM3) (2,2';6',2''-terpyridine-KM1, kM1', kM1''), (OC-6-a)-, bis[hexafluorophosphate(1-)] [9C1] (CA INDEX NAMS)

CM 1

CRN 478818-93-8 CMF C38 H29 N9 O Ru CCI CCS

16919-18-9 P6 P CCS

478819-02-2 CAPLUS
Ruthenium(2+), aqua(N-ethyldipyrido[3,2-a:2*,3*-c]phenazine-11-carboxamide-kN4, kN5) (2,2*:6*,2**-terpyridine- kN1, kN1*, kN 1**1-, (0C-6-43)-, bis/hexafluorophosphate(1-)] (9CI) (CA INDEX NAME)

N 478819-77-1 CAPLUS
N Ruthenium[2-), (acetonitrile) (N-[2-[2-[2-(2,1-di)dycxypropoxy)ethoxy]

CM 1

CRN 478819-76-0 CMF C45 H43 N9 O6 Ru CCI CCS

PAGE 1-A

PAGE 1-B

CM 2

CRN 16919-18-9 CMF F6 P CCI CCS

RN 478819-80-6 CAPLUS
CN Ruthenium(2+), aqua(N-[2-(2-(2,3-dihydroxypropoxy)ethoxy]etho

PAGE 1-A
OH
|
HO-CH2-CH-CH2-O-CH2-CH2-O-CH2-CH2-NH-C

PAGE 1-B

RN 478415-64-4 CAPLUS
CN Dipyrido[3,2-s:2',3'-c]phenazine-11-carboxamide, N-[11-[bis(4-methoxyphenyl)]phenylmethoxyl-14,14-bis(4-methoxyphenyl)-14-phenyl-3,6,9,13-tetraoxatetradec-1-yl]- (9CI) (CA INDEX NAME)

PAGE 1-A

1'')-, (OC-6-43)-, bis(hexafluorophosphate(1-)) (9CI) (CA INDEX NAME)

CM 1

CRN 478619-79-3

CMP C43 H42 N8 07 Ru

CCI CCS

PAGE 1-B

PAGE 1-A

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> CM 2 CRN 16919-18-9 CMP F6 P CCI CCS

F- F- F-

478415-63-3P 478415-64-4P 478819-41-9P
478819-51-1P 478819-57-7P
RL: RCT (Reactant): SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
(DNA-Ru(tpy) (dpp) (CH3CN) | 2 conjugates and their photo crosslinking
studies with complementary DNA strand shows enhanced thermal and
nuclease stability)
478415-63-3 CAPLUS
Dipyrido(3,2-s:2',3'-c)phenazine-11-carboxamide, N-[2-[2-[2-(2,3-dihydroxypropoxy)ethoxy]etho

PAGE 1-B

RN 478819-41-9 CAPLUS

Knthenium(1+), chloro (N-[2-[2-[2-(2,3-dihydroxypropoxy)ethoxy]ethoxy]ethoy]

Jdipyrido(3,2-s-2',3'-c]phenazine-11-carboxamide
kN4, kN5(2,2':6',2''-terpyridine-kN1, kN1', kN

1'')-, (Oc-6-43)-, hexafluorophosphate(1-) (9C1) (CA INDEX NAME)

CM 1

CRN 478819-40-8

CMF C43 H40 C1 N8 O6 Ru

CCI CCS

PAGE 1-A

PAGE 1-B

CH 2

478819-51-1 CAPLUS
Ruthenium(1+), chloro [N-{11-hydroxy-14,14-bis(4-methoxyphenyl)-14-phenyl-3,6,9,13-terreoxatetradec-1-yl]dipyrido[3,2-e:2',3'-c]phenazine-11-carboxamide-xk4, xk5](2,2':6',2''-terpyridine-k11, xk1', xk1'')-, (OC-6-4)-, hexafluorophosphate(1-)(9C1) (CA INDEX NAME)

CM 1

CRN 478819-50-0 CMP C64 H58 C1 N8 O8 Ru CCI CCB

PAGE 1-A

PAGE 1-R

DOCUMENT NUMBER: TITLE:

AUTHOR(S):

137:257239
SAR by MS: A Ligand Based Technique for Drug Lead
Discovery Against Structured RNA Targets
Svayze, Eric E.; Jefferson, Elizabeth A.;
Sannes-Lowery, Kristin A.; Blyn, Lawrence B.; Risen,
Lisa M.; Arakawa, Satoshi; Osgood, Stephen A.;
Hofstadler, Steven A.; Griffey, Richard H.
Ibis Therapeutics, A Division of Isis Pharmaceuticals
Inc., Carlebad, CA, 92008, USA
Journal of Medicinal Chemistry (2002), 45(18),
3161-3181

CORPORATE SOURCE:

SOURCE:

Journal of Medicinal Chemistry (2002), 45(18),
3316-3619
CODEN: JMCMAR; ISSN: 0022-2623
DUCLMENT TYPE: American Chemical Society
Journal
LANGUAGE: Snjieh
TOTHER SOUNCE(S): CASRRACT 137:257239
AB A technique for lead discovery vs. RRA targets utilizing mass spectrometry
(MS) screening methods is described. The structure-activity relationships
(GAR) derived from assaying weak binding motifs allows the pharmacophores
discovered to be elaborated via "SAR by MS" to higher affinity ligands.
Application of this strategy to a subdomain of the 335 FRNN afforded a new
class of compds. with functional activity.

IT EL: CST (Combinatorial study, 2012-28sified); PAC (Pharmacological
ectivity); TRU (Theraputic use); BIOL (Biological study); CMBI
(Combinatorial study); USES (Uses)
(SAR by MS: ligand-based technique for drug lead discovery against
structured RNA targets)

RN 42119-54-6 CAPLUS

CN 6-Quinoxalinecarboxamide, 1,2,3,4-tetrahydro-2,3-dioxo-N-propyl- (9CI)
(CA INDEX NAMS)

462119-55-7 CAPLUS 6-Quinoxalinecarboxs (CA INDEX NAME) examide, 1,2,3,4-tetrahydro-2,3-dioxo-N-2-propenyl- (9CI)

462119-56-8 CAPLUS 6-Quinoxalinecarboxamide, 1,2,3,4-tetrahydro-2,3-dioxo-N-(phenylmethyl)-(SCI) (CA INDEX NAME)

CRN 16919-18-9 CMF F6 P CCI CCB

478819-57-7 CAPLUS
Ruthenium, chloro(mono(2-(bis(4-methoxyphenyl)phenylmethoxy)-1-[12-(dipyrido(3,2-a:2',3'-clphenezin-11-yl- kM4, kM5)-12-oxo-2,5,8-trioxa-11-ezadodec-1-yl]ethyl] butanedioato(2,2':&',2''-terpyridine-kM1, kM1', kM1')-, (OC-6-43)- (SC) (CA INDEX MAME)

PAGE 1-A

PAGE 1-B

REFERENCE COUNT:

THERE ARE 71 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

71 L13 ANSWER 45 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2002:558411 CAPLUS

462119-69-3P
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); TRU (Therapeutic use); BIOL (Biological atudy); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (SAR by MS: ligand-based technique for drug lead discovery sgainst structured RNA targets) 462119-69-3 CAPLUS 6-Quinoxalinecarboxamide, N-[3-[5-[[(1R)-3-amino-1-(1-piperaxin/learboxyll-propyllamino]carboxyll-2-(uranyl]propyl]-1,2,3,4-tetrahydro-2,3-dioxo- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

38 THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 46 OF 181 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: AUTHOR(S):

CAPLUS COPYRIGHT 1006 ACS on STN
1002:501576 CAPLUS
137:208794
Electron-deficient columnar plestic crystals
Book, Harald; Babeau, Annick; Seguy, Isabelle;
Jolinat, Pascale; Destruel, Pierre
Centre de Recherche Paul Pascal, CNRS, Pessac, 33600,
Fr.

CORPORATE SOURCE:

Fr.
ChemPhysChem (2002), 3(6), 532-535
CODEN: CPCHFT; ISSN: 1439-4235
Wiley-VCH Verlag GmbH
Journal

PUBLISHER:

DOCUMENT TYPE: LANGUAGE:

ISHER: Miley-VCH Verlag GmbH
MINT TYPE: Journal
UNGE: Journal
The compound 5,6,11,12,17,18-hexaexatrinaphthylene-2,8,15-tricarboxylic acid
(I) was synthesized from hexaketocyclohexane octahydrate and
3,4-diaminohenzoic acid which were refluxed in glacial acetic acid. The
synthesis of the corresponding seters of I (R = Et. Pr. Bu. n-penty),
n-hexyl, n-heptyl, n-octyl, n-nonyl, 1-ethylpropyl, 2-methylpropyl,
2-ethylbutyl, 2-tehoxyethyl, 2-isopropoxyethyl, rac-3-methoxybutyl, and
ditethoxymethyl)methyl is also described. Furthermore, the compound
3,5,3',5'',5'',5''',5'''-hexapentyloxycarbonyl-2,4,6-triphenyl-1,3,5triezine was prepared by transesterification of the hexamethyl homolog which
was refluxed with K2CO3, 1-pentanol, and 1-bromopentane. The compose were
cheracterized by KRD, MS, H-NGR spectroscopy, DSC, optical-absorption
spectroscopy, and cyclic voltammetry. They form columnar mesophases with
a widely differing degree of order going from highly fluidic nematic via
hexagonal liquid crystallinity to plastic crystallinity. The electron

transport in the strongly electron-deficient compound I is discussed.
444579-19-59
RL: PEP (Physical, engineering or chemical process); PRP (Properties); PYP
(Physical process); SFN (Synthetic preparation); PREP (Preparation); PROC
(Process)
(preparation and electronic and optical properties of)
444579-19-5 CAPUIS
Diquinoxalino[2,3-s:2',3'-c]phenazine-2,8,15-tricarboxylic acid, tributyl
ester (9CI) (CA INDEX NAME)

444579-17-1P 444579-18-4P 444579-20-8P
444579-21-9P 444579-22-DP 444579-23-1P
444579-21-7P 444579-22-DP 444579-23-1P
444579-21-7P 444579-23-PP
444579-20-0P 444579-21-PP
444579-20-0P 444579-21-1P
RL: PEP (Physical, engineering or chemical process); PRP (Propertiee); PYP
(Physical process); SPN (Synthetic preparation); PREP (Preparation); PROC
(Process)
(preparation and liquid-crystalline transition temps. of)
444579-17-3 CAPUS
Diguinoxalino[2,3-s:2',3'-c]phenszine-2,8,15-tricarboxylic acid, triethyl
ester (9CI) (CA INDEX NAME)

444579-18-4 CAPLUS
Diquinoxalino[3,3-a:2',3'-c]phenazine-2,8,15-tricarboxylic acid, tripropyl eacer (9CI) (CA INDEX NAME)

444579-20-8 CAPLUS Diquinoxalino [2,3-e:2',3'-c]phenazine-2,8,15-tricarboxylic acid, tripentyl ester (921) (CA INDEX NAME)

444579-21-9 CAPLUS
Diquinoxalino(2,3-a:2',3'-c]phenazine-2,8,15-tricarboxylic acid, trihexyl
eater (9C1) (CA INDEX NAME)

444579-22-0 CAPLUS
Diquinoxalino{2,3-a:2',3'-c|phenazine-2,8,15-tricarboxylic acid, triheptyl
ester (9CI) (CA INDEX NAME)

444579-23-1 CAPLUS
Diquinoxalino[2,3-a:2',3'-c]phenazina-2,8,15-tricarboxylic acid, trioctyleater (9CI) (CA INDEX NAME)

444579-24-2 CAPLUS
Diquinoxaline(2,3-a:2',3'-c]phenazine-2,8,15-tricarboxylic acid, trinonyl ester (9CI) (CA INDEX NAME)

444579-25-3 CAPLUS
Diquinoxalino[2,3-a:2',3'-c]phenaxine-2,8,15-tricarboxylic acid,
tris(1-ethylpropyl) ester (9CI) (CA INDEX NAME)

444579-26-4 CAPLUS
Diquinoxalino[2,3-a:2',3'-c]phenezine-2,8,15-tricarboxylic acid,
tris(1-methylpropyl) ester (SCI) (CA INDEX NAME)

444579-27-5 CAPLUS Diquinoxalino(3,3-e12',3'-c)phenazine-2,8,15-tricerboxylic acid, tria(2-ethylbutyl) ester (9CI) (CA INDEX NAMS)

444579-30-0 CAPLUS
Diquinoxalino(2,3-e:2',3'-c)phenazine-2,8,15-tricarboxylic acid,
tria(3-methoxybutyl) ester (9CI) (CA INDEX NAME)

444579-31-1 CAPLUS
Diquinoxalino(2,3-e:2',3'-c)phenazine-2,8,15-tricarboxylic acid,
tris(2-ethoxy-1-(ethoxymethyl)ethyl) ester (9CI) (CA INDEX NAME)

444579-28-6 CAPLUS
Diquinoxalino[2,3-e:2',3'-c]phenazine-2,8,15-tricarboxylic ecid,
tris(2-ethoxyethyl) ester (9CI) (CA INDEX NAME)

444579-29-7 CAPLUS
Diquinoxalino[3,3-a:2',3'-c]phenezine-2,8,15-tricerboxylic acid,
trie[2-(1-methylethoxylethyl] ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

REPERENCE COUNT:

26 THERE ARE 26 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSMER 47 OP 181 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1002.481308 CAPPUS

117.194994

AUTHOR(S):

AUTHOR(S):

AUTHOR(S):

AUTHOR(S):

AUTHOR SOURCE:

CORPORATE SOURCE:

I lie Therapeutice, A Division of Isia Pharmaceuticals Inc. Carlabed, CA, 29008. USA.

CORPORATE SOURCE:

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PAGE 1-C

RN 452338-60-2 CAPLUS
CN 6-Quinoxalinecarboxamide, N-[2-[[(35,65,98)-9-amino-3-[3[(aminoininemethyl)amino]propyl]-6-[(4-aminophenyl)methyl]3,4,5,6,7,8,9,10,11,12-decahydro-5,8,12-trioxo-2H-1,4,7,11benzoxatriazacyclotetradecin-14-yl]amino]-2-oxoethyl]-1,2,3,4-tetrahydro-Nmethyl-1,3-dioxo-[9C1] (CA INDEX NAND)

Absolute stereochemistry.

PAGE 1-C

PAGE 1-A

RN 452338-61-3 CAPLUS
CN 6-Quinoxalinecarboxamide, N-[3-[[(15,65,95)-9-amino-3-[3[(aminoiainomethy)]amino]propyl]-6-[(4-aminophenyl]methyl]3,4,5,6,7,8,9,10,11,12-decahydro-5,8,12-trioxo-2k-1,4,7,11benzoxatrizazeycoltetradecin-14-yl]amino]-3-oxopropyl]-1,2,3,4-tetrahydro2,3-dioxo- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

PAGE 1-C

452338-62-4 CAPLUS
6-Quinoxalinecarboxamids, N-(4-{{(38,68,98)-9-amino-3-[3-{(aminoishnomethyl)amino]propyl}-6-{(4-aminophenyl)amthyl]-3,4,5,6,7,8,9,10,11,12-decahydro-5,8,12-trioxo-3H-1,4,7,11-bensoxatriasacyclotatradecin-14-yl]amino]-4-oxobutyl]-1,2,3,4-tetrahydro-2,3-dioxo-(9C1) (OA INDEX NAME)

Absolute stereochemistry.

PAGE 1-C

RN 452338-63-5 CAPLUS
CN 6-Quinoxalinecarboxamide, N-[6-[[[38,68,98]-9-amino-3-[3-[[aminoiminomethy]] amino] propyl]-6-[[4-aminophenyl]methyl]3,4,5,6,7,8,9,10,11,12-decahydro-5,8,12-trioxo-2H-1,4,7,11-benzoxatrizazeycolotetradecin-14-yl]amino]-6-oxohexyl]-1,2,3,4-tetrahydro2,3-dioxo- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-C

REFERENCE COUNT: THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 48 OF 181 ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE: INVENTOR(S):

CAPLUS COPYRIGHT 2006 ACS on STN
2002:184970 CAPLUS
136:221460
Improvements relating to water treatment
Walker, Gavin Michael
The Queen's University of Belfast, UK
PCT Int. Appl., 17 pp.
CODEN: PIXXD2
Patent PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

PAGE 2-A

NH₂

REFERENCE COUNT: THERE ARE 5 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2006 ACS on STN 2002:157741 CAPLUS 136:200190 Benzimidazoles and analogues and their use as L13 ANSWER 49 OF 181 ACCESSION NUMBER:

DOCUMENT NUMBER:

TITLE:

Benzimidazoles and analogues and their use as neutrophil inhibitors Bush, Rodney Dean; Hershberger, Paul Mitchell; Young, Judith Anne; Kasibhatla, Bhavani The Procter & Gamble Company, USA PCT Int. Appl., 56 pp. CODRN: PIXXD2 Patent English INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

OTHER SOURCE(S):

| 1 | PA1 | ENT I | NO. | | | KIN | D | DATE | | | APPL | ICAT | ION | NO. | | D | ATE | |
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| 1 | WO | 2002 | 0163 | 27 | | A1 | | 2002 | 0228 | | WO 2 | 001- | US25 | 224 | | 2 | 0010 | 810 |
| 1 | WO | 2002 | 0163 | 27 | | Cı | | 2002 | 0801 | | | | | | | | | |
| | | W: | AE. | AG. | AL. | AM, | AT. | AU. | AZ. | BA. | BB. | BG. | BR. | BY. | BZ. | CA. | CH. | CN. |
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MARPAT 136:200190

401791-58-0P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (intermediate; benzimidazoles and analogs and use as neutrophil

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO. DATE 20020314 APPLICATION NO. WO 2001-GB3994 DATE 20010906 MO 2002001512 Al 20020114 WO 2001-081994 20010906
M: AE, AO, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, ER, ES, FI, GB, GD, GR, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, KK, MM, MM, MX, MZ, DO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SI, TJ, TM, TT, TZ, LQ, LOU, US, UZ, VN, YU, ZA, ZM, AM, AZ, BY, KO, KZ, MD, RU, TJ, TM, RM: GH, GM, KE, LS, MM, MZ, SD, SI, SZ, TZ, UQ, ZW, AT, BE, CM, CV, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BP, BJ, CF, CG, CI, CM, GA, GM, GO, GM, ML, MR, NE, SD, TD, TO AU 2001086652 AS 20010212 AU 2001-86052 20010906 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
US 2004020659 Al 20040205 US 2001-36153 20010606 R: LT, LI, LU, NL, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
US 2004020659 Al 20040205 US 2001-36163 20010606 AD 20040205 US 2001-36163 20010607 AD 20040205 US 2001-36160 20040205 US 2001-361600205 US 2001-36160 20010506 US 2001-36160 20010506 US 2001-361600205 US 2001-3616

206058-73-3 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-{[[4-[[(2,3-dichloro-6-quinoxaliny]]carbonyl]amino]sethyl]-3-sulfophenyl]sethyllamino]-9,10-dixor., disodium salt (9CI) (CA INDEX NAME)

PAGE 1-A

Title compds. I [X, Y * heteroatoms wherein at least X or Y is (un) substituted nitrogen; Z * C, two C atoms or a heteroatom; R - alkyl, aromatic ring, carbocyclic aliphatic ring, halo, haloalkyl, heteroatkyl, heterocyclyl, H, OH, NH2, SH, OCH3; R1-3 - alkyl, aromatic, carbocyclic aliphatic, halo(alkyl), heteroalkyl, heterocyclyc aliphatic, halo(alkyl), heteroalkyl, heterocyclyc aliphatic, H; L = C:O-A-NR3, NR3-A-C:O, RNN-A-C:O-A-NR3, SH2, aromatic ring, carbocyclic aliphatic ring, haloalkyl, heteroalkyl, heterocyclyl, H; B = alkyl, aromatic ring, carbocyclic aliphatic ring, carbocyclic aliphatic ring, end that linke R4-5 into a cyclic ring structure; if G is nil, R4 = alkyl-carboxy, aryl-carboxy, etc.; R5 = H, alkyl, aromatic ring, carbocyclic aliphatic ring, carbocyclic aliphatic ring, carbocyclic aliphatic ring, carbocyclic aliphatic ring, halo, haloalkyl, heteroalkyl, heteroalkyl, heteroalkyl, pato, haloalkyl, heteroalkyl, lower heteroalkyl, etc.] were propered For instance, 5-benzimidaxolacarboxylic acid was coupled to L-phenylalanine benzyl ester (DMF, EDAC, NOSE, ELIN) and the resulting aside debenzylated (MeON, H3-Pd/C) to give II. Compds. I are useful for the treatment and prevention of diseases and conditions aschemic-reperfusion injury.:

10:791-70-69

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PRSP (Preparation); USES (Uses)

(Grug; benzinidexoles and analogs and use as neutrophil inhibitors)
401791-70-6 CAPLUS

L-Phenylalanine, N-(6-quinoxalinylcarbonyl)- (GCI) (CA INDEX NAME)

L-Phenylalanine, N-(6-quinoxalinylcarbonyl)-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry

REFERENCE COUNT

THERE ARE 11 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 50 OF 181 CAPLUS COPYRIGHT 2006 ACE on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
136:151182
136:151182
Antimicrobial biaryl compounds
Jefferson, Blizabeth Ann; Swayze, Eric
1sis Pharmaceuticale, Inc., USA
PCT Int. Appl., 44 pp.
CODEN; PIXD2

DOCUMENT TYPE:
PARENT
ENGISER

DOCUMENT TYPE: LANGUAGE: English 2

FAMILY ACC. NUM. COUNT:

| AIE | W1 . | INFOR | WAT I | ON: | | | | | | | | | | | | | | |
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| | WO | 2002 | 0096 | 48 | | A2 | | 2002 | 0207 | | WO : | 2001- | US24 | 067 | | 2 | 0010 | 801 |
| | WO | 2002 | 0096 | 48 | | A3 | | 2002 | 0627 | | | | | | | | | |
| | _ | W: | AB. | AG. | AL. | AM. | AT. | AU. | AZ. | BA. | 88. | BG. | BR. | BY. | BZ. | CA. | CH. | CN. |
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| | EP | | | | | | | | | | | 2001- | | | | | | |
| | | R: | AT, | BĖ, | CH, | DE, | DK, | ES, | FR, | GB, | GR, | IT, | LI, | LU, | NL, | SE, | MC, | PT, |
| | | | | | | | | | | | | TR | | | | | | |
| | JP | 2004 | 5194 | 21 | | T2 | | 2004 | 0702 | | JP 2 | 2002- | 5152 | 03 | | 2 | 0010 | 801 |
| RIOF | RIT | APP | LN. | INFO | . : | | | | | | US 2 | -000 | 6301 | 22 | | A 2 | 0000 | 801 |
| | | | | | | | | | | | WO 2 | 1001- | US24 | 067 | 1 | W 2 | 0010 | 801 |
| THEF | R SC | URCE | (8): | | | MAR | PAT | 136: | 1511 | 62 | | | | | | | | |
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PATENT INFORMATION:

| PATENT | NO. | KIND DA | TE | APPLIC | ATION NO. | DATE |
|--------------|-------------|-----------|----------|----------|------------|-----------------|
| | | | | | | |
| US 6323 | 227 | B1 20 | 011127 | US 199 | 9-259528 | 19990226 |
| US 6080 | 767 | A 20 | 0000627 | US 199 | 7-884405 | 19970627 |
| WO 9900 | 356 | A1 19 | 990107 | WO 199 | 8-US13550 | 19980626 |
| W: | AL, AM, AT, | AU, AZ, B | A, BB, B | G, BR, B | Y, CA, CN, | CU, CZ, DE, DK, |
| | EE, ES, FI. | GB, GE, G | H, HU, I | L. 18. J | P, KE, KG, | KP, KR, KZ, LC, |
| | LK, LR, LS, | LT, LU, L | V, MD, M | G, MK, M | N, MW, MX, | NO, NZ, PL, PT, |
| | RO, RU, SD, | SE, SG, S | I, SK, S | L, TJ, T | M, TR, TT, | UA, UG, US, UZ, |
| | VN, YU, ZW | | | | | |
| RW: | GH, GM, KE, | LB. MW. S | D. SZ. U | G, ZW, A | T. BE. CH. | CY, DE, DK, ES, |
| | FI, FR, GB, | GR, IE, I | T, LU, M | C, NL, P | r, se, br, | BJ, CF, CG, CI, |
| | CM, GA, GN, | ML, MR, N | E, SN. T | D. TG | | |
| PRIORITY APP | LN. INFO.: | | | US 199 | 6-9485P | P 19960102 |
| | | | | US 199 | 7-884405 | A2 19970627 |
| | | | | WO 199 | 8-US13550 | A1 19980626 |
| | | | | WO 199 | 6-0920770 | A2 19961223 |
| OTHER SOURCE | (s): | MARPAT 13 | 6:5913 | | | |

Title compds. I [R = H, OH, NH2; Rl = R2 = H; or RlR2 = :NH9; R3 = H, COJR6, COR6, CON(R6)2, CHJOR7, CHJSR7; R4 = H, alkyl, alkyl-Q, thioheterocyclyl, (CHJCH2)Ar, (CH:CH)AR. CHJAr; R5 = alk(en/yn)yl, cycloalk(en/yl, heterocyc)(en/yl, aryl, heteroaryl, fused systems, etc.; R6 = H, lower alkyl; R7 = H, lower alkyl, aralkyl, lower acyl, aroyl, heteroaroyl; R8 = H, lower alkyl; R9 = H, R100CC, R10C, CH, Cycnon, R10CO, OHC, lower alkyl, OJN, Y1'Y2'H; R10 = alkyl, aralkyl, heteroaralkyl; Y1', Y2' = H, alkyl; O = TO, R75, Y1Y3H; Y1, Y2' = H, alkyl, aryl, aralkyl; or one of Y1 and Y2 = acyl or aroyl and the other is as given; Ar = aryl or heteroaryl; n = 0-2] and their pharmaceutically acceptable salts, prodrugs, N-oxides, hydrates, and solvetes are useful as Factor Xa inhibitors. For example, 4-(pyridin-3-yl)benzic acid was amidated with tert-Bu 3-eminopropionate-Hcl via the acid chloride, and the resulting B-acylamino ester underwent a sequence of (1 o-alkylation with 5-iodo-2-[(2-methoxyethoxy)methoxy]benzyl bromide, (2) acidic deprotection

Biaryls I [X = CH, O, S, N, NH; Y = CH, N; n = 0, 1; one of R1 and R2 = (un) substituted CONRH2, COONH3, CRIVALS, SOONNA; and the other is H or R3; one of R5 and R6 = NICOR7, NHSGOR7, NHSGOR7 and the other is H, R4; O = mino acid or peptide residue; R3 = H, halogen, (un) substituted NH2, NHCOR7, R4 = H, halogen, hydroxyl, amino, carboxyl, alkyl, alkynyl, 5-16 member carbocycle or heterocycle were prepd for use as antimicrobial agents. Thus, polymer-supported piperszine was acylated with 5-bromo-2-thiophenecerboxylic acid, coupled with 3-RNCSHBS(OH2, and acylated with 2.3-dioxobenzopyraxine-6-carboxylic acid to give the biaryl II. In a coupled bacterial transcription-translation assay II had an ICSo of 25 µM.
395648-38-1P
RL: BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(Uses)
(preparation of acylaminobiarylcarboxamides as bactericides)
195648-18-1 CAPLUS
6-Quinoxalinecarboxamide, 1,2,3,4-tetrahydro-2,3-dioxo-N-[2-oxo-2-[[3-[5-(1-piperazinylcarbonyl)-2-thienyl]phenyl]amino]ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\$$

L13 ANSMER 51 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2001:863510 CAPLUS
DOCUMENT NUMBER: 136:5913
TITLE: Preparation of substituted N-[(aminoiminomethyl or aminomethyl) propyl amides as Factor Xa inhibitors
RINVENTOR(S): Klein, Scott I.; Guertin, Kevin R.; Spada, Alfred P.;
PATENT ASSIGNEE(S): Aventie Pharmaceuticals Products Inc., USA CODEN: USXXAM
DOCUMENT TYPE: Pauls, R.; Cont.-in-part of U.S. Ser. No. 884,405.

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT:

of the MEM group, and conversion to the Me eater, (3) Pd-catalyzed cyenation of the iodide, and (4) Pinner reaction and ammonolysis of the nitrile, to give title compound II. Three example compds showed Ki values of 13.0-94.0 nM in a Factor Xe assay, 46 nM to 1.72 µM in a trypsin assay, and 477 nM to 2.71 µM in a thrombin assay.

219673-02-6P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therspeutic use); BIOL (Biological atudy); PREP (Preparation); UBES (Uses) (preparation of substituted [(aminominomethyl)- or [(aminomethyl)-propyl propyl amidce as Factor Xe inhibitors) 219673-03-6 CAPIUS
Benzenepropanoic acid, 3-(aminominomethyl)- a-[1-[(6-quinoxalinylcarbonyl)aminolethyl)-, methyl ester (SCI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 52 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN

2001:587526 CAPLUS 135:318478

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

135:318478
Traceless, Self-Cleaving Solid- and Solution-Phase
Parallel Synthesis of 3,4,7-Trisubstituted
3,4-Dihydroquinoxalin-2-ones
Laborde, Edgardo; Peterson, Brian T.; Robinson, Louise
Telik Inc., South San Francisco, CA, 94080, USA
Journal of Combinatorial Chemistry (2001), 3(6), AUTHOR(S): CORPORATE SOURCE: SOURCE:

strong acids or to maintain an inert atmospheric, thereby preserving the chiral integrity of the starting \(\alpha\) amino acid and facilitating the generation of libraries in a high-throughput parallel format.

167941-10-49
RL: SPN (Synthetic preparation). PREP (Preparation).

(traceless self-cleaving solid- and solution-phase parallel synthesis of 3.4.7-trisubstituted 3.4-dihydroquinoxalin-2-ones)

167941-10-4 CAPPUS
6-Quinoxalinecarboxamide, N-[13-fluorophenyl]methyl]-3.4-dihydro-2-(1-sethylethyl)-3-oxo- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 53 OF 181 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

AUTHOR (S): CORPORATE SOURCE:

CAPLUS COPYRIGHT 2006 ACS on STN
2001:465005 CAPLUS
135:319476
Cold pad-batch dyeing of Lyocell
Siedow, K.
DyStar Textilfarben OmbH y Co., Frankfurt am Main,
Germany
Revieta de Quimica Textil (2001), 151, 42, 44-46,
48-53
CODEN: ROTEDJ; ISSN: 0300-3418
Asociacion Espanola de Quimicos y Coloristas Textiles
Journal
Spanish

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

MAGE: Spanish
Methods to reduce crease and rub marks on Lyocall fabrics and suitable
cold pad-batch dyeing processes, i.e., sods and silicate, are described.
Finishing methods and reagents for woven fabrics with a peach skin effect
(fleece) and with a smooth (non-fibrillated) surface are outlined.
Operation conditions for cold pad-batch dyeing of Lyocall were determined and
dyed Lyocall was compared to viscose fabrics in terms of color intensity,
dye affinity, tailing test, fixing yield, washing-out, etc. Types of cold
pad-batch process include soda process for Levafix dyes, water glass
methods for Remazol dyes; recommendations are given for dye selection for
each type. methods for Remaxol dyes; recommendations are given to, up each type.

206058-73-1, Levafix Brilliant Blue 8-B
RL: NUI Other use, unclessified; USES (Uses)
(dye selection and perameters for cold pad-batch dyeing and finishing of Lyocell)

206058-73-3 CAPJUS
2-Anthracenesulfonic acid, 1-amino-4-[[4-[[(2,3-dichloro-6-quinoxalinyl]carbonyl]amino]methyl]-3-sulfophemyl]methyl]amino]-9,10-dinydro-9,10-dioxo-, disedium salt (9CI) (CA INDEX NAME)

of the (Ru2.-ODN)-DNA duplexes is found to increase considerably (ATM = 12.8-31.4*), depending upon the site of the covalent attachment of the tethered (Ru(phen) ddpp2)2. complex, or the chirality of the [Ru(phen) 2dpp2)2.-linker tethered at the siddle of the ODN. compared to the Unlabeled counterpart. Gross differences in CD between the [Ru(phen) 2dpp2)2.-linker tethered and the siddle of the ODN. compared to the unlabeled counterpart. Gross differences in CD between the [Ru(phen) 2dpp2]2.-tethered and the native DNA duplexes showed that the global duplex conformation of the former has considerably altered from the B-type, but is still recognized by DNase I. The thermal melting studies, CD measuremente, as well as DNase I digestion data, are interpreted as a result of intercalation of the dpps molety, which is realized by threading of the Ru(phen)2 complex part through the DNA duplex core. DNase I footprinting with four disasterementically pure middle ([Ru(phen) 2dpp2)2.-DON). DNA duplexes furthermore showed that the tethered [Ru(phen) 2dpp2]2.-Inker chirality dictates the stereochem. accessibility of various phosphodiester moleties (around the intercalation site) toward the cleavage reaction by the enzyme. The disasterementically pure ruthenium-modified duplexes, with the well-defined m-stack, will be useful to explore stereochem. dependent energy- and electron-transfer chemical to understand oxidative demagnet to the DNA double helix as well as the long-range energy- and electron-transfer processes with DNA as a 142906-42-7P 342906-43-8P 342906-45-0P 342906-45-0P 342906-45-1P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthesis of (Ru(phen) 3dpp2)2--tethered oligodeoxyribonucleotides) 342906-42-7 CAPLUS

Ruthenium(2+), [N-[11-hydroxy-14, 14-bis (4-methoxyphenyl)-14-phenyl-3, 4, 9, 9, 13-tetraeoxatetradec-1-y] dipyrido(3, 2-a:2', 3'-c) phenazine-11-carboxanide-kNA, kNS)bis (1, 10-phenanthroline-kN1, kN10)- (OC-6-33)- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

●2 Na

ACCESSION NUMBER:

ACCESSION NUMBER:

DOCUMENT NUMBER:

TITLE:

Synthesis of [Ru(phen)2dppz]2*-tethered oligo-DNA and studies on the metallointercalation mode into the DNA duplex

AUTHOR(S):

CORPORATE SOURCE:

Department of Biooryanic Chemistry Biomedical Center, University of Uppala, Uppala, Swed.

Journal of the American Chemical Society (2001), 133(15), 3551-3562

PUBLISHER:

DOCUMENT TYPE:

JOURNAL JACSAT; ISSN: 0002-7863

American Chemical Society

JOURNAL JACSAT; JOURNAL JACSAT; JOURNAL JACSAT; JOURNAL JACSAT; JOURNAL JACSAT; JOURNAL

PAGE 1-B

342906-43-8 CAPLUS
Ruthenium[2+], [1-{[bis(4-methoxyphenyl)phenylmethoxy]methyl]-13(dipyrido[3,2-a;2',3'-c]phenazin-11-yl- kN4, kN5)-13-oxo-3,6,9trioxa-12-azatridec-1-yl P-(2-cyanoethyl)-N,N-bis(1methylethyl)phosphonamidite]bis(1,10-phenanthroline- kN1, kN10)-,
(OC-6-33)- (9C1) (CA INDSK NAME)

PAGE 1-A

PAGE 1-B

342996-45-0 CAPLUS
Ruthenium(1-), [mono(1-{[bis(4-methoxyphenyl)phenylmethoxy]methyl}-13(dipyrido(3,2-s;2',3'-c]phenazin-11-yl- «N4, «N5)-13-oxo-3,6.9trioxa-12-exetridec-1-yl] butanedioacolbis(1,10-phenanthrolineKN, «N10-), (OC-6-33) (9C1) (CA INDEX NONE)

PAGE 1-B

342906-46-1 CAPLUS
Ruthenium(2+), [N-{2-(2-[2-(2,3-dihydroxypropoxy)ethoxy]ethoxy]ethoxy]ethoyl]dipyrido[3,2-2:3,3'-c]phenazine-11-carboxamide- kN4,kN5]bis(1,10-phenanthroline- kN1,kN10)-, (OC-6-33)- (9CI) (CA INDEX NAME)

DAGE 1-A

321835-65-8 CAPLUS
Ruthenium(2+), [N-[2-oxo-2-(tricyclo[3.3.1.13,7]dec-2ylaminolethyl]dipyrido[3,2-a:2*,3*-c]phenazine-11-carboxamideKM4, KN5]bis(1,10-phenanthroline-KM1, KN10)-,
(OC-6-13)-(SCI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 7 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LI3 ANSWER S6 OF ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: AUTHOR(S): CORPORATE SOURCE:

ANSWER 56 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
SSION NUMBER: 2000:739213 CAPLUS
NEWLY NUMBER: 14:72834
E: Lyocell: cold pad-batch dyeing process. Part 2
OK(E): Siedow, K.
ORATE SOURCE: DyStar Textilfabren CMSH & Co., Prankfurt Am Main,

SOURCE.

Germany Tinctoria (2000), 97(8), 31-36 CODEN: TINCAW; ISSN: 0040-7984 Edizioni Ariminum

PUBLISHER:

DOCUMENT TYPE: LANGUAGE: AB Operation

INHER: Editioni Ariminum

MENT TYPE: Journal

UAGE: Italian

Operation conditions for cold pad-batch dyeing of Lyocell were determined and
dyed Lyocell was compared to viscose fabrics in terms of color intensity,
dye affinity, tailing test, fixing yield, washing-out, etc. Types of cold
pad-batch process include soda process for Levafix dyes, water glass
methods for Remazol dyes; recommendations are given for dye selection for
each type.

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THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 55 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 2001:64269 CAPLUS

DOCUMENT NUMBER: TITLE:

134:128208
134:128208
Detection of biomolecules by sensitizer-linked substrates for biomolecules
Gray, Harry B.; Crane, Brian R.; Winkler, Jay R.;
Dmochowski, Ivan Julian; Milker, Jonathan J.; Dunn,
Alexander Robert
California Institute of Technology, USA
PCT Int. Appl., 174 pp.
CODEN: PIXXD2
Patent
English

PATENT ASSIGNER(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.

quinoxalinyl)carbonyl]amino]methyl]-3-sulfophenyl]methyl]amino]-9,10-dihydro-9,10-dioxo-, disodium salt (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

REFERENCE COUNT: THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ACCESSION NUMBER:
DOCUMENT NUMBER:
DOCUMENT NUMBER:
DIMERISATIONS AVAILABLE IN THE RE FORMAT

AUTHOR(S):

AUTHOR(S):

AUTHOR(S):

AUTHOR SOURCE:

CORPORATE SOURCE:

Dimerisation of Cibacron Blue F JGA and other dyes:
influence of salts and temperature
Maria; Fisichella, Salvatore
Dipertisento di Scienze Chimichs, Universita di
Catania, Catania, 59125, Italy
Dyes and Pigments (2000), 46(3), 129-137

CODEN: DYPIDX: ISSN: 0143-7208
Elseviers Science Ltd.
DOCUMENT TYPS:

Journal
LANGUAGE:

Biglief

AB The monomer-dimer equilibrium of Cibacron Blue F JGA (CB) and five other dyes
(Levafix Brilliant Blue EB, Reactive Scarlet 017, Methyl orange, Basic
Blue 3, and Chicego Blue Skyl have been investigated in water and in the
presence of Kalapo4. Aggregation of CB has been aloe examined in the

on non-linear least-square fitting procedure was applied, it was found that the diserization consts. depend on the extension of organic mols. and the number of sulfonic groups. In the case of CB, cations had a greater effect on the equilibrium than anions. Anal. of the calculated spectra for

mer and dimer of Basic Blue 3 after deconvolution allowed us to specify the geometry of the dimer. 206058-73-3, Levafix Brilliant Blue BB RL: PRP (Properties); TBM (Technical or engineered material use); USBS

(Uses)

(influence of salts and temperature on aqueous associative dimerization of

s)
206058-73-3 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[[4-{[(2,3-dichloro-6-quinoxaliny])carbony]amino]methyl]-3-sulfophenyl]methyl]amino]-9,10-dihydro-9,10-dioxo-, diaodium malt (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

●2 Na

REFERENCE COUNT:

THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 58 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2000:700553 CAPLUS DOCUMENT NUMBER: TITLE:

134:4913 Synthesis of some sulfonamide derivatives with potential antibacterial activity

El-Din, Nabaweya Sharaf Faculty of Pharmacy, University of Tanta, Egypt Chemistry of Heterocyclic Compounds (New York) (Translation of Khimiya Geteroteiklicheskikh Soodinenii) (2000), 36(4), 449-454 CODEN: CHCCAL; ISSN: 0009-3122 Consultants Bureau AUTHOR (S): CORPORATE SOURCE: PURI.I SHED

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S):

UNENT TYPE: Journal

DUAGET

STATE

Brish

English

SMAGE

SMOURCE(S): CASERACT 134.4913

Some new quinoxaline-6-sulfonanide and phhalazine-6-sulfonanide derivs.

were synthesized in 61-684 yields by treating the corresponding
quinoxaline- and phhalazine-6-sulfonyl chlorides with the appropriate
samine (PrNN2, 1-smino-2-propanol, glycine, p-H2NCSH4CO2H, morpholine,
piperarine). The smjority of the prepared compda. showed antibacterial
activity.

11217-24-0P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); SPN (Synthetic preparation); BIOL (Biological
study, unclassified); SPN (Synthetic preparation); BIOL (Biological
study); PREP (Preparation)
(preparation and antibacterial activity of)

112170-26-0 CAPLUS

Olycine, N-(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]- (9CI)
(CA INDEX NAME)

REFERENCE COUNT: THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 36

LI3 ANSWER 59 OP 161
ACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE:
INVENTOR(S):
INVENTOR(S):
SOURCE:
LAVELUS AVENTAGE AND CALIFORM ACS ON STN
(arcylamino)alkancates and analogs as factor Xa inhibitors
INVENTOR(S):
SOURCE:
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PATENT INFORMATION

| PAT | TENT | NO. | | | KIN | D | DATE | | | APPL | ICAT | ION | NO. | | D | ATE | |
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| US | 6080 | 767 | | | A | | 2000 | 0627 | | US 1 | 997- | 8844 | 05 | | 1 | 9970 | 627 |
| WO | 9724 | 118 | | | A1 | | 1997 | 0710 | | WO 1 | 996- | US20 | 770 | | 1 | 9961 | 223 |
| | W: | AL, | AM, | AT, | AU, | AZ, | BB, | BG, | BR, | BY, | CA, | CH, | CN, | CZ, | DE, | DK, | EE, |
| | | ES. | PI. | GB, | GE. | KU, | IL. | IS, | JP, | KE, | KG, | KP, | KR, | KZ, | LK, | LR, | LS, |
| | | LT, | LU, | LV, | MD, | MG, | MX, | MIN, | MW, | MX, | NO, | NZ, | PL, | PT, | RO, | RU, | БD, |
| | | SB, | SG, | SI, | SK, | TJ, | TM, | TR, | TT, | UA, | UG, | us, | UZ, | VN | | | |
| | RW: | KE, | LS. | MW, | SD, | SZ, | υa, | AT, | BE, | CH, | DE, | DK. | ES, | PI. | FR, | GB, | GR, |
| | | IE, | IT, | LU, | MC, | NL, | PT, | SE, | BF, | BJ, | CF, | CG, | CI, | CM, | GA, | GN, | ML, |
| | | MR, | NE, | SN, | TD, | TG | | | | | | | | | | | |
| CA | 2264 | 556 | | | AA | | 1999 | 0107 | , | CA 1 | 998- | 2264 | 556 | | 1 | 9960 | 626 |
| | | | | | | | | | | | | | | | | | |

| WO | 99003 | 156 | | | A1 | | 1999 | 0107 | | WO | 1998 - | US13 | 550 | | | 19980 | 626 |
|----------|---------|------|------|-----|------|-----|------|------|-----|----|--------|-------|-----|-----|------|--------|-----|
| | W: | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BF | , BY, | CA, | CN, | CU, | cz | , DE, | DK, |
| | | EE, | ES, | PI. | GB, | GE, | GH, | HU, | IL, | 18 | , JP, | KE, | KO, | KP, | KR | , KZ, | LC, |
| | | LK, | LR, | LS, | LT, | LU, | LV, | MD, | MG, | MX | , MN, | MW, | MX, | NO, | NZ | PL, | PT, |
| | | RO, | RU, | SD, | SE, | SG. | SI, | SK, | SL. | TJ | , TM, | TR, | TT. | UA, | UG | . US, | UZ, |
| | | VN, | YU, | 2W | | | | | | | | | | | | | |
| | RW: | GH, | GM, | KE, | LS, | MW, | SD, | SZ, | UG, | Z¥ | , AT, | BE, | CH, | CY, | DE | , DK, | ES, |
| | | FI, | FR, | GB, | GR, | IE, | IT, | LU, | MC, | NI | , PT, | SE, | BF, | BJ, | CF | , co, | CI, |
| | | CM, | GA, | GN, | ML, | MR, | NE, | SN, | TD, | TG | | | | | | | |
| UA | 98817 | 771 | | | A1 | | 1999 | 0119 | | ΑU | 1998- | 6177 | 1 | | | 19980 | 626 |
| UA | 74117 | 73 | | | B2 | | 2001 | 1122 | | | | | | | | | |
| EP | 93106 | 0 | | | A1 | | 1999 | 0728 | | EP | 1998- | 9317 | 26 | | | 19980 | 626 |
| | R: | AT, | BE. | CH, | DE, | DK, | ES, | FR, | GB, | GR | , IT, | LI, | LU, | NL, | SE | , MC, | PT, |
| | | IE, | | | RO | | | | | | | | | | | | |
| | 98060 | | | | | | | | | | 1998- | | | | | | |
| | 20015 | | | | | | | | | | | | | | | | |
| AP | 1061 | | | | Α | | 2002 | 0424 | | AP | 1999- | 1467 | | | | 19980 | 626 |
| | | | | KE, | LS. | MW, | SD, | sz, | UG, | 2W | 1 | | | | | | |
| ZA | 98056 | 64 | | | A | | 1999 | 113 | | ZA | 1998- | 5664 | | | | 19980 | 629 |
| NO | 99008 | 54 | | | A | | 1999 | 0423 | | Ю | 1999- | 854 | | | | 19990: | 223 |
| | 31479 | | | | | | 2003 | | | | | | | | | | |
| | 63232 | | | | | | | | | | | | | | | 19990: | 226 |
| | 62778 | | | | B1 | | 2001 | 0821 | | | 1999- | | | | | 19990 | |
| PRIORITY | / APPI | N. | NFO. | . : | | | | | | US | 1996- | 9485 | P | - 1 | ₽ : | 19960 | 102 |
| | | | | | | | | | | WO | 1996- | US20 | 770 | | A2 : | 19961 | 223 |
| | | | | | | | | | | | 1997- | | | | | 19970 | |
| | | | | | | | | | | | 1998- | | | | | | |
| | | | | | | | | | | WO | 1998- | US13: | 550 | 1 | | 19980 | 626 |
| OTHER SC | OURCE (| (s): | | | MARI | TAS | 133: | 7386 | L | | | | | | | | |

H2NCRIR2ZCH2CHRICHRINGSCORS [R1, R2 = H; R1R2 = NR9; R3 = H, COR6, CO2R6, CON1R6]2,CH2OR7, CH2GR7; R4 = H, (hydroxy) alkyl, aminoalkyl, (CM2CH2)nR, (CH:CH)nR, CM2R7 = (un) substituted (hetero) arryl, R5 = (ar) alk(en)yl, heterocyclyl, (hetero) arryl, etc.; R6, R8 = H or alkyl; R7 = H, alkyl, acyl, (heterolaryl, etc.; R9 = H, OR, alkoxy (carbonyl), alkanoyl, etc.; Z = phenylene; n = 0-2] were prepared as factor Xa inhibitors (no data). Thus, 4-(NC) CSHCHICHCHCHC) were cyclocondepsed with 4-(MHO) CSHAN:CHCH.(CMPh [preparation each given) to give, after N-deprotection, β-lactam I. The latter was N-acylated by 4-Phc6H2COL and the product hydrolyzed to give, after amination/esterification, title compound II.

219673-02-6P
RJ: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); TMU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of u-amidinobenzyl-\$-(aroylamino) alkanoates and analogs as factor Xa inhibitors)
219673-02-6 CAPLUS
Benzenepropanoic acid, 3-(aminoiminomethyl)- u-{1-{(6-quinoxalinylcarbonyl)amino|ethyl]-, methyl ester (9CI) (CA INDEX NAME)

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE PORMAT

REFERENCE COUNT:

L13 ANSNER 60 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
13114414
Synthesis and Assembly of Self-Complementary Cavitands
AUTHOR(S):
CORPORATE SOURCE:
Self-Complementary Cavitands
Renslo, Adam R: Tucci, Pabio C:; Rudkevich, Dmitry
M: Rebek, Julius, Jr.
Skeggs Institute for Chemical Biology and the
Department of Chemistry, Scripps Research Institute,
La Jolla, CA, 2027, USA
Journal of the American Chemical Society (2000),
122(19), 4573-4552
CODEN: JACSAT; ISSN: 0002-7863
American Chemical Society
DOCUMENT TYPE:
JOURNELL Begins
Begins
Begins

DOCUMENT TYPE: LANGUAGE: English

NAMN TYPE: Journal NUMBER: Smglish Cavitands with self-complementary shapes were prepared by the covalent attachment of admantane guest mole. to the upper rim of the host structures. Relatives of the "melf-folding" cavitands, these new structures possess a seam of intramol. hydrogen bonds that stabilize the folded conformation. Their self-complementary shapes result in the formation of noncovalent dimer; of considerable kinetic and thermodn. stability (-AG995 - 4.5-6.5 kcal/mol in p-xylene-dio). The dimerization of the cavitands is reversible and subject to control by selvent and temperature The dimerization process is enthalpically favored and entropy opposed and occurs with significant enthalpy-entropy compensation. 109239-81-8P 242129-41-5P 270553-44-5P RL: RCT (Reactant): SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant): printegs and inclusion dimerization of self-complementary cavitands bearing admantyl recognition sites) 108229-81-8 CAPLUS (CA INDEX NAME)

242129-41-5 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(tricyclo{3.3.1.13,7]dec-1-ylmethyl)- (9CI) (CA INDEX NAME)

270563-44-5 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, tricyclo[3.3.1.13,7]dec-1-ylaethyl ester (9C1) (CA INDEX NAME)

270563-41-2P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
(hydrogenolysis; synthesis and inclusion dimerization of
self-complementery cavitands bearing adamantyl recognition sites)
270563-41-2 CAPLUS
13.23:14.22-Dimetheno-15H, 17H, 19H, 21H-benzo(2',3')benzo(2'',3'')[1,7]benzo
dioxonino(3''',2''':9'',10'')[1,4]benzodioxonino(6',5'';9',10')[1,4]benzod
dioxonino(6',5''9,10][1,4]benzodioxonino(2,3-b]quinoxaline-27-carboxylic
acid, 2,3,9,10,5,5,6-bexakis([1-oxoccyt])aminol-15,17,19,21-tetraundecyl-,
phenylmethyl ester, (15R,178,19R,218)-rel- (9CI) (CA INDEX NAME)

PAGE 1-C

— (CH₂)₆-Me

Me- (CH2) 6-

PAGE 1-B

PAGE 3-B

(СН₂) 10 - ме

270563-45-6P
RL: PEP (Physical, engineering or chemical process); PRP (Properties); SPN (Synthetic preparation); PREP (Preparation); PROC (Process) (no dimerization; synthesis and inclusion dimerization of self-complementary cavitands bearing adamantyl recognition sites) 270563-45-6 CAPLUS
6-Quinoxalinecarboxylic acid, 2-chloro-3-[[(24R,26R,28R,34S)-31-hydroxy-2,3,9,10,16,17-hexakis[(1-oxocctyl)amino|-24,26,28,34-tetraundcyl-22,30-methano-24H,26H,28H-tetraundcyl-22,30-methano-24H,26H,28H-tetraundcyl-22,30-methano-24H,26H,28H-tetraundcyl-22,30-methano-24H,26H,28H-tetraundcyl-27,30-methano-24H,26H

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

IT 242143-99-3P
RL: PEP (Physical, engineering or chemical process); PRP (Properties); SPN
(Synthetic preparation); PREP (Preparation); PROC (Process)
(prepared and characterized in present paper and mis-assigned in earlier
paper; synthesis and inclusion dimerization of self-complementary
cavitands bearing admantyl recognition sites)

RN 242143-99-3 CADLUS
CN 13,23:14,22-01metheno-15H,17H,19H,21H-benzo[2',3']benzo[2'',3''][1,7]benzo
dioxonino[3'',2'':9'',10''][1,4]benzodioxonino[6'',5'':9',10''][1,4]benzodioxonino[6'',5'':9',10'][1,4]benzodioxonino[2,3-b]quinoxaline-27-carboxamide,
2,3,9,10,35,36-beaksis[(1-oxoctyl)amino]-N-tricyclo[3,3.1,13,7]dec1ylmethyl)-15,17,19,21-tetraundecyl-, stereoisomer (9CI) (CA INDEX NAME)

PAGE 1-A

Me- (CH2) 6-

PAGE 1-C

- (CH₂)₆-Me

PAGE 2-A

RL: PEP (Physical, engineering or chemical process): PRP (Properties): SPN (Synthetic preparation): PRSP (Preparation): PROC (Process) (synthesis and inclusion diserization of self-complementary cavitands bearing ademantyl recognition sites)
270561-64-9 CAPLUS

270563-64-9 CAPLUS
6-Quinoxal inecerboxamide, 2-chloro-3-[(31-hydroxy-2,3,9,10,16,17-hexakis[(1-oxoocty1)amino]-24,26,28,34-tetraundecy1-22,30-methano-244,262,284-tetraundecy1-22,30-methano-244,262,284-tetraundecy1-22,30-methano-244,262,284-tetraundecy1-22,35-methano-244,262,284-tetraundecy1-22,15,6]'lbis[1,4]benzodioxonin-21-yl]oxy]-N-(tricyclo[3.3.1.13,7]dec-1-ylmethyl)-, stereoisomer (9C1) (CA INDEX NAME)

STRUCTURE DIAGRAM IS NOT AVAILABLE ***
270563-65-0 CAPLUS
6-Quinoxalinecarboxylic acid, 2-chloro-3-[[31-hydroxy-2,3,9,10,16,17-hexakls[[1-oxoocty]]amino]-24,26,28,34-tetraundecyl-22,30-methano-244,26H,28H-tetrabenzo[b,b',e,e'][1,7]benzodioxonino[3,2-j:5,6-j']bis[1,4]benzodioxonino11-yl]boxyl-, tricyclo[3,3,1.13,7]dec-1-ylmethylester, stereoisomer (9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
REFERENCE COUNT: 37 THERE ARE 3

THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ACCESSION NUMBER:
DOCUMENT NUMBER:
1171LE:
AUTHOR(S):
AUTHOR(S):
CORPORATE SOURCE:
CORPORATE SOURCE:

SOURCE:

PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
DOCUMENT TYPE:
LANGUAGE:
BECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

2000:122112 CAPLUS
120:20766
Sequence-Recognition and Cleavage of DNA by a
Netropsin-phenezine-di-N-oxide Conjugate
Helissey, Philippe; Olorgi-Rensult, Sylviene; Colson,
Plerre; Houssier, Claude; Bailly, Christian
Laboratoire de Chinke Therapeutique Faculte des
Sciences Pharmaceutiques et Biologiques, UMR
CKRS-Universite Rene Descartes no. 8638, Paris, 75270,
Fr.

SOURCE:
Bioconjugate Chemistry (2000), 11(2), 219-227
CODEN: BCCHES; ISSN: 1043-1802
American Chemical Society
Journal
LANGUAGE:
English

SIDECHINGS: SCRIEG: ISSN: 1043-1802

PUBLISHER: American Chemiscal Society

Journal

LANGUADE: DAN-BERGERS: ISSN: 1043-1802

American Chemical Society

Journal

LANGUADE: American Chemical Society

Journal

LANGUADE: American Chemical Society

American Chemical Ch

PAGE 3-B

(CH₂)₁₀ - Mo

270561-88-7
RL: PRU (Pormation, unclassified); PRP (Properties); FORM (Formation, nonpreparative)
(synthesis and inclusion dimerization of self-complementery cevitands bearing admentyl recognition sites)
270561-88-7 CAPLUS
6-Quinoxalinecarboxamide, 2-chloro-1-[[11-hydroxy-2,3,9,10,16,17-hexakis[(1-cxoxoctyl)amino]-24,26,28,34-tetraundecyl-22,30-methano-24H,26H,28H-tetrabenzo[b,b',e,e'][1,7]benzodioxonino(3,2-):5,6[']bis[1,4]benzodioxonin-21-ylloxy]-N-tricyclo[3,3.1.13,7]dec-1-yl-, stereoisomer, compd. with stereoisomer of 2-chloro-3-[[31-hydroxy-2,3,9,10,16,17-hexakis[(1-cxoxoctyl)amino]-24,26,8,34-tetraundecyl-22,30-methano-24H,26H,28H-tetraundecyl-121,30-methano-24H,26H,38H-tetraundecyl-121,30-methano-24H,26H,38H-tetraundecyl-121,30-methano-24H,36H,34H-121,30-methano-24H,36H,34H-121,30-methano-24H,36H,34H-121,30-methano-24H,36H,34H-121,30-methano-24H,36H,34H-121,30-methano-24H,36H,34H-121,30-methano-24H,36H,34H-121,30-methano-24H,36H,34H-121,30-met

CM 1

CRN 270563-64-9 CMF C158 H228 Cl N9 O15

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

CM 2

CRN 270563-63-8 CMF C157 H226 Cl N9 O15

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
IT 270563-64-9P 270563-65-0P

atudy, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (R 128; sequence-recognition and cleavege of DNA by netropsin-phenazine-di-N-oxide conjugate R-128) 260416-9-9 (APLUS

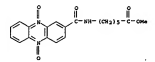
26916-09-9 CAPLUS
2-Phenezinecarboxamide, N-[6-[[5-[[[5-[[(3-emino-3-ininopropy])-mlno]carbonyl]-1-methyl-1H-pyrrol-3-yl]amino]carbonyl]-1-methyl-1H-pyrrol-3-yl]amino]carbonyl]-1-methyl-1H-pyrrol-3-yl]amino]-6-oxohexyl]-, 5,10-dioxide, monohydrochloride (9CI) (CA INDEX NAME)

• HC1

PAGE 1-B

260389-77-19 260389-78-49 260389-79-59
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
(sequence-recognition and cleavage of DNA by netropsin-phenazine-di-Noxide conjugate R-128)
260389-77-3 CAPLUS
Hexanoic acid. 6-[(2-phenazinylcarbonyl)amino]-, methyl ester (9CI) (CA
INDEX NAME)

260189-78-4 CAPLUS
Rexanoic acid, 6-[([\$,10-dioxido-2-phenazinyl)carbonyl]amino]-, methyl
ester [901] (CA INDEX NAME)



9-5 CAPLUS
acid, 6-[[(5,10-dioxido-2-phenazinyl)carbonyl]amino]- (9CI) (CA

THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 62 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN

L13 ANSWER 62 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2000:41754 CAPLUS
112:208087
TITLE: 2000:41754 CAPLUS
112:208087
TITLE: 2000:41754 CAPLUS
112:208087
TITLE: 2000:41754 CAPLUS
112:208087
Dipyrido[3,2-a:2*,3*-c]phenazine-tethered oligo-DNA:
synthesis and thermal stability of their DNA:
DNA and DNA: NNA duplexes and DNA: DNA:
AUTHOR(S): 20sipov, Dimitri; Zamaratski, Edouard; Chattopadhyaya,
Jyoti
CORPORATE SOURCE: Department of Bioorganic Chemistry, Biomedical Center,
University of Uppsala, Swed.
Helvetica Chimica Acta (1993), 82(12), 2186-2200
COURN: HCACAV; ISSN: 0018-019X
Verlag Helvetica Chimica Acta
DOCUMENT TYPS: Journal
LANGUAGE: Splish
AB Dipyrido[3,2-a:2*,3*-c]phenazine (dppz) derivs. were conjugated to 3-mer
and 18-mer DNA (ODN) at a site without nucleobase, either at the 5*- or
3*-end or at a internucleotide position, via linkers of 7, 12, or 18 atoms
lengths. These dpp: linked ODNs were synthesized using novel backbone
glycerol phosphoramidities: glycerol, serving as artificial nucleoside
without nucleobase, was modified to amines which were suitable for the
subsequent key reaction with dpp-carboxylic acid. The products of these
reactions were then transformed to the standard phosphoramidite derive. or
used for loading on a CPO support. The dpps-modified ODNs were
subsequently assembled in the usual manner using automated solid-phase DNA
synthesis. The 3-mer ODN-dpps conjugates were tested for their ability to
form stable duplexes with target DNA or RNA strands (D11 or R11) while to
18-mer ODN-dpps conjugates were tested for their ability to
conjugated dpps derivative increases the stability of DNA or DNA and DNA
- RNA duplexes, typically by a ATm of 7.3-10.99 and
4.5-7.4*, resp., when the dpps is tethered at the 5*- or

PAGE 1-B

259796-27-5 CAPLUS
Dipyrido[3,2-a:2',3'-c]phenazine-11-carboxamide, N-[17-hydroxy-20,20-bis[4-methoxypheny1)-20-phenyl-3,6,9,12,15,19-hexaoxaeicos-1-yl]- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

259796-37-7 CAPLUS
Dipyrido(3,2-a:2',3'-c)phenazine-11-carboxamide, N-[3-(2,3-dihydroxypropoxy)propyl]- (9CI) (CA INDEX NAME)

the 5'- or 3'-end, with a ATm varying from 3.8-11.1°. The insertion of a dppz building block at the center of a 9-mer results in a considerably poorer stability of the corresponding DNA - DNA duplexes (ATm = 0.5 to 4.2°) and DNA - RNA duplexes (ATm = 1.5 to 0.3°), while the replacement of one interior nucleotide by a dppz building unit in the corresponding 8-mer ODN does not reveal the formation of any duplex at all. Different types of modifications in the middle of the 18-mer ODN, in general, do not lead to any triplex formation, except when the dppz derivative is tethered to the ODN through a 12-atom-long linker.
259784-25-19 25978-64-64 2 25978-40-2DP, CPG bound 25978-64-3-DP, CPG bound 259786-43-DP, CPG bound 559786-43-DP, CPG bound 7 C

bound

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthesis and thermal stability of dipyrido[phenazine]-tethered oligo-DNA and their DNA/RNA duplexes and triplexes)

259796-25-3 CAPUUS

Dipyrido[3,2-a:2',3'-c]phenazine-11-carboxamide, N-[3-[42,2-dimethyl-1,3-dioxolan-4-yl)methoxy]propyl]- (9CI) (CA INDEX NAME)

259796-26-4 CAPLUS
Dipyrido(3,2-a:2',3'-c)phenazine-11-carboxamide, N-[11-hydroxy-14,14-bis(4-methoxyphenyl)-14-phenyl-3,6,9,13-tetraoxatetradec-1-yl]- (CA INDEX NAME)

PAGE 1-A

259796-38-8 CAPLUS
Dipyrido[1,2-a:2',3'-c]phenazine-11-carboxamide, N-[3-[3-[bis(4-mathoxyphenyl)phenylmethoxy]-2-hydroxypropoxy]propyl]- (9CI) (CA INDEX NAME)

PAGE 1-B

PAGE 1-A

PAGE 1-A

259796-40-2 CAPLUS Butanedicic acid, sono[1-[[bis(4-methoxyphenyl)phenylmethoxy]methyl]-2-[3-(dipyrido[3,2-a:2',7-e]phenazin-11-ylcarbonyl)amino]propoxy]ethyl] ester (9c1) (CA INDEX NAMS)

- C- CH2- CH2- CO2H

259796-43-5 CAPLUS
Butanedioic acid, mono[1-{[bis(4-methoxyphenyl)phenylmethoxy]methyl}-13-dipyrido(3,2-a:2',3'-c]phenazin-11-yl-13-oxo-3,6,9-trioxa-12-azatridec-1-yl) ester (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

259796-44-6 CAPLUS
Butanedioic acid, mono[1-[[bis(4-methoxyphenyl)phenylmethoxy]methyl]-19-dipyrido[3,2-a:2',3'-c]phenazin-11-yl-19-oxo-3,6,9,12,15-pentaoxa-18-azanonadec-1-yl] ester [9CI) (CA INDEX NAMS)

PAGE 1-A

259796-40-2 CAPLUS
Butanedioic acid, mono[1-{[bis(4-methoxyphenyl)phenylmethoxy]methyl]-2-[3-[dipyrido],2-a:2',3'-c]phenazin-11-ylcarbonyl)amino]propoxy]ethyl] ester (9C1) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

259796-24-2P 259796-39-9P 259796-40-2P 259796-41-1P 259796-41-1P 259796-42-4P 259796-43-5P 259796-46-8P 259796-46-8P 259796-47-9P RI: SPN (Synthetic preparation); PREP (Preparation) (synthesis and thermal stability of dipyrido(phenaxine)-tethered oligo-DNA and their DNA/RNA duplexes and triplexes) 259796-24-2 CAPLUS Dipyrido(3.2-a:2',3'-c)phenaxine-11-carboxamide, N-(2-hydroxyethyl)-N-methyl- (9CI) (CA INDEX NAME)

259796-39-9 CAPLUS
Phosphoranidous acid, bis(1-methylethyl)-, 1-[(bis(4-methoxylethyl)phenyl)phenylmethoxylmethyl)-2-(3-[(dipyrido[3,2-a:2',3'-c]phenazin-11-ylcarbonyl)amino]propoxylethyl 2-cyanoethyl ester (9CI) (CA INDEX NAME)

PAGE 1-B

PAGE 1-A

PAGE 1-B

259796-42-4 CAPLUS
Phosphoremidous acid, bis(1-methylethyl)-, 1-{[bis(4-methyylphenyl)phenylmethoxylmethyl]-19-dipyrido[3,2-si2',3'-c]phenazin-11-yl-19-oxo-1,6,9,12,15-pentaoxa-16-azanonadec-1-yl 2-cyanoethyl ester (9CI) (CA INDEX NAME)

PAGE 1-B

RN 259796-41-5 CAPLUS
CN Sutanedioic acid, mono[1-[[bis(4-methoxyphenyl)phenylmethoxy]methyl]-13dipyrido[3,2-a:2',3'-c]phenagin-11-yl-13-oxo-3,6,9-trioxa-12-azatridec-1yl] ester (9CI) (CA INDEX NAME)

PAGE 1-A

RN 259796-44-6 CAPLUS
CN Butanddioic exid, mono[1-[|bis(4-methoxyphenyl)phenylmethoxy]methyl]-19dipyrido[3,2-a;2',3'-c]phenazin-11-yl-19-oxo-3,6,9,12,15-pentaoxa-18azanonadec-1-yl] ester (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

RN 259796-45-7 CAPLUS
CN Thymidine, thymidylyl-(3' -5')-2'-deoxycytidylyl-(3' -5')-2'-deoxycytidylyl-(3' -5')-2'-deoxycytidylyl-(3' -5')-2'-deoxyadenylyloxy[2-[[3-[(dipyrido[3,2-a:2',3'-c]phenszin-11-ylcarbonyl)amino[propoxy]methyl]-1,2-ethanediyl]oxyphosphinico-(3' -5')-2'-deoxyadenylyl-(3' -5')-2'-deoxycytidylyl-(3' -5')-(CA INDEX NAME)

Absolute stereochemistry.

нο,

PAGE 1-B

PAGE 2-B

RN 259796-46-8 CAPLUS

CN Thymidine, thymidylyl-(3'-s')-2'-deoxycytidylyl-(3'-5')-2'deoxycytidylyl-(3'-5')-2'-deoxyadenylyloxy(2-(12-dipyrido[3,2a:2',3'-c]phenazin-11-yl-12-oxo-2,5,8-trioxa-11-azadodec-1-yl)-1,2ethanediylloxyphosphinico-(3'-s')-2'-deoxyadenylyl-(3'-s')2'-deoxycytidylyl-(3'-s')-2'-deoxyadenylyl-(3'-s')(CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

N 259796-47-9 CAPLUS
N Thymidine, thymidylyl-(3' +5')-2'-deoxycytidylyl-(3' +5')-2'-deoxycytidylyl-(3' +5')-2'-deoxycytidylyl-(3' +5')-2'-deoxyadenylyloxy[2-(18-dipyrido[3,2-a:2',3'-c]phenazin-11-yl-18-oxo-2,5,8,11,14-pentaoxa-17-azaoctadec-1-yl)-1,2-ethanediyl loxyphosphinico-(3' +5')-2'-deoxyadenylyl-(3' +5')-2'-deoxyctidylyl-(3' +5')-2'-deoxyadenylyl-(3' +5')-(5CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 3-C

PAGE 4-A

thioformamide, formanide acetal or thioformamide acetal, in the presence of a halogenating agent. Examples of suitable halogenating agents include but are not limited to thionyl chloride, phosepne, and phosepne derives. Reactants containing more than one addni. N-C-C-N group may also be used to prepare compds, with two or more inidezolium groups, by the procedures of the present invention. Certain compds. of the invention prepared from reactants with multiple N-C-C-N groups may have both unreacted N-C-C-N moieties and substituted imidezolium groups. N. g., 1,2-bis(dimethylamino)-2,3-dimethyl-6-(2-pyridylamino)-1; imm perchlorate (90%) was prepared by reaction of 7,8-dimethyl-2,3-di-2-pyridylaminosaline with Ne2NCSM/SOCI2, followed by treatment with aqueous NaCIO4.

246518-08-1P 246518-09-2P 246518-10-5P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

246518-09-2 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-di-2-pyridinyl-, octyl ester (9CI) (CA INDEX NAME)

246518-10-5 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-di-2-pyridinyl-, decyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS

REFERENCE COUNT: THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSMER 63 OF 181 CAPLUS COPYRIGHT 2006 ACS on STM
ACCESSION NUMBER: 1599:719212 CAPLUS
1132:64246
Synthesis of some sulfonamide derivatives with potential entibacterial activity
AUTHOR(S): E1-Din, Nebaweya Sharaf
CORPORATE SOURCE: Department of Pharmaceutical Chemistry, Paculty of Pharmacy, University of Tanta, Tanta, Egypt
OTIENTAL OCURRING, 158N: 0970-0200.

PUBLISHER: OTIENTAL SCIENTIFIC COURS. OTIENTAL SCIENTIFIC C

were synthesized. Most of the products showed antibacterial activity, 112170-26-09
RI: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
(preparation and antibacterial activity of)
112170-26-0 CAPLUS
Glycine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]- (9CI)
(CA INDEX NAME)

THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 64 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1999:670141 CAPLUS DOCUMENT NUMBER: 131:286517

Imidazolium cations and processes for their

Imidazolium cations and processes for their preparation Donovan, Robert J.; Morgan, Robert J. The Rockefellar University, USA U.S., 23 pp., Cont.-in-part of U.S. 5,874,587. CODEN: USXXAM Patent INVENTOR (8): PATENT ASSIGNEE (S): SOURCE:

DOCUMENT TYPE: LANGUAGE: glish

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

TITLE:

PATENT NO. APPLICATION NO. DATE KIND DATE US 5969150 US 5874587 PRIORITY APPLN. INFO.: OTHER SOURCE(S):

US 5565150 A 19991019 US 1996-124546 19980729 US 5574517 A 19990223 US 1996-673687 19960625 RITY APPLN. INFO.:

RSOURCE(S): CASREACT 131:286517; MARPAT 131:286517 The present invention relates to novel imidazolium compds. and improved processes for the preparation of imidazolium cations with one or more imidazolium moieties optionally substituted with the same or different substituents, which are prepared from a reactant with at least one N-C-C-N group, by reacting with with an N-substituted or N,N-disubstituted

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LI3 ANSWER 65 OF 181
ACCESSION NUMBER:
DOCUMENT NUMBER:
1399:62554 CAPLUS
131:346148
131:346148
Dimeric analogues of non-cationic tricyclic aromatic carboxamides are a new class of cytotoxic agents Spicer, Julie A.; Gemage, Swarna A.; Attell, Oraham J.; Pinlay, Graeme J.; Baguley, Bruce C.; Denny, William A.

CORPORATE SOURCE:
Auckland Cancer Society Research Centre, Paculty of Medicine and Health Science, The University of Auckland, Auckland, N. Z.

SOURCE: Anti-Cancer Drug Design (1999), 14(3), 281-289

PUBLISHER: OXford University Press
DOCUMENT TYPE:

Auckland, Auckland, N. Z.

Anti-Cancer Drug Design (1999), 14(3), 281-289

CODEN: ACDDEA; ISSN: 0266-536

PUBLISHER: Oxford University Press

Journal

LANGUAGE: Begins

As a series of tricyclic arcmatic carboxamides, and their corresponding dimeric

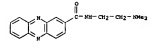
analoge, were prepared and their growth-inhibitory properties were evaluated
in a series of cell lines. The dimeric compde, were prepared by reaction of
the appropriate acids with carbonyl-1,1'-dimidazole, isolating the
resulting imidaxolides, and reacting these with a stoichiometric amount of
the appropriate acids with carbonyl-1,1'-dimidazole, isolating the
resulting imidaxolides, and reacting these with a stoichiometric amount of
the differing inhibitory potencies, with the known
nitronephthalimide (mitonafide) and acridine-4-carboxamide (DACA) being
the most potent. The corresponding bis analoge, linked by a
(CH2)3NMM (CH2)3 chain, were generally more potent, with the largest
increases (dimer/monomer ratio 20- to 30-fold) seen for the
nitronaphthalimides and the phenazines. Based on the intrinsic
cytotoxicity of the monomers and the highest degree of increase in
cytotoxicity of dimerization, the most interesting chromophores appear to
be the acridine-4-carboxamide and phenazine-1-carboxamide. Both of these
compds. showed significant growth delays (.apprx.6 days) in an in vivo
colon 35 tumor model in mice.

IT 25064-07-22 (APLUS
R) (Shological activity or effector, except adverse); BSU (Biological
study, unclassified); SPN (Synthetic preparation); TNU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); USES (Usee)
(dimeric analogs of non-cationic tricyclic aromatic carboxamides are a new
class of cytotoxic agents in relation to structure)

2064-07-2 (APLUS
2-Phenazinecarboxamide, N,N'-[(methylimino)di-3,1-propanediyl]bis(CA INDEX NAME)

250684-04-9
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(Uses)
(dimeric analogs of non-cationic tricyclic aromatic carboxamides are a new class of cytotoxic agents in relation to structure)
250684-04-9 CAPLUS
2-Phenarinecarboxamide, N-[2-(dimethylamino)ethyl]- (9CI) (CA INDEX NAME)



REFERENCE COUNT:

L13 ANSMER 66 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1999;455125 CAPLUS
DOUMEMENT NUMBER: 131:1993190
TITLE: Self-Complementary Cavitands
AUTHOR(8): Sension Adam R.; Rudkevich, Dmitry M.; Rebek, Julius

AUTHOR(S):

Renelo, Adam R.; Rudkevich, Dmitry M.; Rebek, Julius Jr.
Skegge Institute for Chemical Biology and The Department of Chemistry, The Scrippe Research Institute, Le Jolla, CA. 92037, USA

SOURCE:

JOURNELL LE Jolla, CA. 92037, USA
PUBLISHER:

DOCUMENT TYPE:
JOURNELL JOURNELL LE JOLLA, CA. 92037, USA
American Chemical Society
JOURNELL LE JOLLA, CA. 92037, USA
Vase-like cavitands with single adamentyl groupe covalently bound on the upper rim via veriable-length espectre were prepared and their inclusion processes studied in competing and noncompeting solvente. JMR and computer modeling suggested that in noncompeting p-xylene the adamentane halle were included quant, within the highly shielding environment of the cavitand binding socket in self-complementary dimeric assemblies. In CDC11 the assembly is much weaker, and variable temperature binding studies provided AB = -10.6 Kcal/mol and AB = -241.5 eu.; e., the binding is enthalpically favorable and entropically unfavorable.

TRI RCT (Reactant); RACT (Reactant or reagent) (coupling with cavitand diol; preparation of cavitands with covalently bound adamantyl groupe on their upper rime and their self-complementary cavity-guest binding in dimeric assemblies)

RN 342125-41-5 CAPLUS
CN 6-00inoxalineacetoxamide, 2,3-dichloro-N-(tricyclo[3.3.1.13,7]dsc-1-ylmethyl)- (SCI) (CA INDEX NAME)

242144-39-4
RL: FMU (Formation, unclassified); PEP (Physical, engineering or chemical process); PRP (Properties); PORM (Formation, nonpreparative); PROC (Process)

(Process)
(preparation of cavitands with covalently bound adamantyl groups on their
upper rims and their self-complementary cavity-guest binding in dimeric
sessemblies)
242144-19-4 CAPLUS
13, 23:14, 22-Dimetheno-15H, 17H, 19H, 21H-benzo [2', 3']benzo [2'', 3''][1,7]benzo
dioxonino[3'', 2''';9'',10''][1,4]benzodioxonino[6'',5'';9',10'][1,4]benzo
dioxonino[6',5':9,10][1,4]benzodioxonino[2,3-b]quinoxaline-27-carboxamide,

CRN 242143-99-3 CMF C158 H227 N9 O15

PAGE 1-A

Me- (CH2)6-

PAGE 1-B

PAGE 1-C

- (CH₂) 6-Me

PAGE 2-A

Me-- (CH₂) 6 - C-

PAGE 2-B Me- (CH2) 10 Me- (CH2) 10 (CH2) 10 - Me

PAGE 3-B

(CH₂)₁₀ – Me

CM 2

CRN 242143-97-1 CMF C157 H225 N9 O15

PAGE 1-A

Me- (CH2)6-

PAGE 1-B

NH-C-(CH₂)6-Me | He-(CH₂)6-C | NH-C-(CH₂)6-Me | C-NH-NH

(СН₂) 10 - ме

PAGE 1-A

Me- (CH2)6-

-- (CH₂)₆-Me

PAGE 2-A

PAGE 1-B

PAGE 1-C

REFERENCE COUNT:

Me- (CH2)10

(CH₂)₁₀-Me THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

PAGE 3-B

(CH₂)₁₀-Me

23

Me- (CH2) 10

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ACCESSION NUMBER:
DOCUMENT NUMBER:
11999:447645 CAPLUS
131:310427
Probing the Role of Polyphenol Oxidation in Mediating Insect-Pathogen Interactions. Galloyl-Derived Electrophilic Traps for the Lymantria disper Nuclear Polyhedrosis Virus Matrix Protein Polyhedrin

AUTHOR(S):

AUTHOR(S):

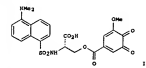
CORPORATE SOURCE:

Foldman, Ken S.: Sambandam, Aruna; Bowers, Katherine E.; Appel, Heidi M.
Department of Chemistry and Pesticide Research Laboratory, The Pennsylvania State University, University Park, PA, 16802, USA
Journal of Organic Chemistry (1999), 64 (16), 5794-5803
CODEN: JOCCAM; ISBN: 0022-1261
AMERICAN:
GI
GI
CASREACT 131:310427

PUBLISHER: DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI

— (СН 3) 6 - ме

PAGE 2-A



- Galloyl-derived orthoquinone probes have been designed, synthesized, and utilized in an ongoing study of insect-pathogen interactions. A stable galloyl-derived orthoquinone O-Me ether modified with both acidic and fluorescent appendages (1) was successful in trapping the model nucleophile cysteine, a test protein bearing a single cysteine residue, and the viral occlusion body matrix protein polyhedrin from Lymantria diepar nuclear polyhedrosis virus (LMNPV), a pathogen of the gypsy moth caterpillar (DMC). This latter observation may be related to the mol. mechanism by which gallotannins decrease LMNPV infectivity in GMC's. Sufficient site isolation was not achieved with a polymer-bound reactive galloyl hydroxysorthoquinone electrophile to permit similar nucleophile trapping to compete with oligomerization.

 245046-12-40 [Synthetic preparation]; PREP (Preparation)
 (trapping cysteine and cysteine-containing proteins by galloyl-derived orthoquinone ather as model for mediating insect-pathogen interactions)

 245048-12-4 [S-(diesthylamino)-1-naphthalenyl]sulfonyl].

 4-mathoxy-2-phenarineoarboxylate (ester) (SCI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: THERE ARE 58 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 68 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1999:147950 CAPLUS
TITLE: 1999:147950 CAPLUS
130:206594
Pyrazine derivatives formed by the reaction of deoxyglucosone with diamino derivatives, antibodies recognizing the product and application in diagnosis
INVENTOR(S): 100:101 diagnosis
Uchida, Yoshiaki; Kurano, Yoshihiro; Ito, Satoru
Pujirebio, Inc., Japan
Ger. Offen. 22 pp.
CODEN: GMXXXX
Patent

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---|-----------------------|------|----------|------------------|----------|
| | | | | | |
| | DB 19837664 | A1 | 19990225 | DE 1998-19837664 | 19980819 |
| | US 6291198 | B1 | 20010918 | US 1998-134388 | 19980814 |
| | JP 11181000 | A2 | 19990706 | JP 1998-249122 | 19980819 |
| | JP 3508563 | B2 | 20040322 | | |
| | GB 2329387 | A1 | 19990324 | GB 1998-18242 | 19980820 |
| ₽ | RIGRITY APPLN. INFO.: | | | JP 1997-240348 A | 19970621 |
| • | т | | | | |

220925-38-1 CAPIDS
6-Quinoxalinecarboxamide, N-[4-[[2-[[1-[2-[4-[5-[(3aS,48,6aR)-hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-y]]-1-oxopenty]]-1-piperaziny]]ethyl]-2,5-dioxo-3-pyrrolidinyl]thiolethyl]amino]-4-oxobutyl]-3-[(2S,3R)-2,3,4-trihydroxybutyl]- (9CI) (CA INDEX NAME)

220928-39-2 CAPLUS 220928-19-2 CAPLUS
6-Quinoxalinecarboxamide, N-[4-[[2-[[1-[2-[4-[5-[(]a8,48,6aR]-hexahydro-2-oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]-1-piperazinyl]ethyl]-2,5-dioxo-3-pyrrolidinyl]thiojathyl]amino]-4-oxobutyl]-2-[(28,3R)-2,3,4-trihydroxybutyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

220882-55-3P 220882-67-7P
RL: RAC (Biological activity or effector, except adverse); BSU (Biological atudy, unclassified); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent) (immunogen, coupling with keyhole limpet hemocyanin for immunization, reaction with BSA to form antigen for antibody isolation; pyraxine derivs. formed by the reaction of deoxyglucosone with diamino derivs.) 220882-55-3 CAPUIS
6-Quinoxalinecarboxamide, N-[4-oxo-4-[(2-(2-pyridinyldithio)ethyl]amino]bu tyl]-3-[(28,3R)-2,3,4-trihydroxybutyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

220882-67-7 CAPLUS
6-Quinoxalinecarboxamide, N-[4-oxp-4-[[2-(2-pyridinyldithio)ethyl]amino]butyll-2-(3,4-trihydroxybutyll- (SCI) (CA INDEX NAME)

Absolute stereochemistry.

L13 ANSWER 69 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NAMBER:
DOCUMENT NUMBER:
130:110161
17TILE: Preparation of substituted N-[(aminoiminomethyl or aminomethyl)] phenyl] propyl amides as Pactor Xa inhibitors
INVENTOR(S): Klein, Scott I.; Guertin, Kevin R.; Spada, Alfred P.; Pauls, Reims W.; Gong, Yong; McGarry, Daniel G. Rhome-Poulene Rorer Pharmaceuticals Inc., USA PCT Int. Appl., 352 pp.
COURCE: PIXXD2
DOCUMENT TYPE: Patent

Patent English

DOCUMENT TYPE: LANGUAGE: FAMILY ACC, NUM. COUNT: PATENT INFORMATION:

| | | | | | | | | | | | | | | | | | DATE | |
|------|------|------|------|------|-----|------|-----|-------|------|-----|----|------|--------|-----|-----|----|----------------|-----|
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| | NO | | | | | | | | | | | | | | | | 19980 | |
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| | | | LK, | LR, | LS, | LT, | LU | , LV, | MD, | MG, | MK | , MN | , MW, | MX, | NO, | NZ | , PL, | PT, |
| | | | RO, | RU, | SD, | SB, | SG | , sı, | εĸ, | SL, | ΤJ | , TM | , TR, | TT, | UA, | UG | , US, | UZ, |
| | | | VN. | YU, | ZW | | | | | | | | | | | | | |
| | | RW: | GH, | GM, | KE, | LS, | MM | , SD, | SZ, | UG, | ZW | , AT | , BE, | CH, | CY, | DE | , DK, | ES, |
| | | | PI, | FR, | GB, | GR, | IE | IT, | LU, | MC, | NL | , PT | , SE, | BF, | ВJ, | CF | , cg, | CI, |
| | | | CM, | GΑ, | GN, | ML, | MR | NE, | SN, | TD, | TG | | | | | | | |
| | บร | 6080 | 767 | | | A | | 2000 | 0627 | | US | 1997 | - 6844 | 05 | | | 19970
19980 | 627 |
| | CA | 2264 | 556 | | | AA | | 1999 | 0107 | | CA | 1998 | -2264 | 556 | | | 19980 | 626 |
| | | | | | | | | | | | AU | 1998 | - 8177 | 1 | | | 19980 | 626 |
| | ΑU | 7411 | 73 | | | B2 | | 2001 | 1122 | | | | | | | | | |
| | EP | | | | | | | | | | | | | | | | 19980 | |
| | | R: | AT, | BE, | CH, | DE, | DK. | ES, | FR, | GB, | GR | , IT | , LI, | LU, | NL, | SE | , MC, | PT, |
| | | | IE, | SI, | | | | | | | | | | | | | | |
| | | 9806 | | | | A | | 1999 | 0831 | | BR | 1998 | -6060 | • | | | 19980 | 626 |
| | J₽ | 2001 | 5005 | 32 | | T2 | | | | | | | | | | | 19980 | |
| | AP | 1061 | | | | A | | | | | | | -1467 | • | | | 19980 | 626 |
| | | ₩: | GH, | GM, | ΚĽ, | LS, | MW, | SD, | SZ. | UG, | ZW | | | | | | | |
| | NO | 9900 | 854 | | | A | | 1999 | 0423 | | NO | 1999 | -854 | | | | 19990 | 223 |
| | NO | 3147 | 58 | | | B1 | | 2003 | 0519 | | | | | | | | | |
| | | | | | | B1 | | 2001 | 1127 | | | | | | | | 19990 | |
| PRIO | RIT | APP | LN. | INFO | . : | | | | | | US | 1997 | -8844 | 05 | | A2 | 19970 | 627 |
| | | | | | | | | | | | US | 1996 | - 9485 | P | | P | 19960 | 102 |
| | | | | | | | | | | - | WO | 1996 | -US20 | 770 | | A2 | 19961 | 223 |
| | | | | | | | | | | 1 | WO | 1998 | -US13 | 550 | 1 | W | 19980 | 626 |
| OTHE | R SC | URCE | (S): | | | MARI | PAT | 130: | 1101 | 51 | | | | | | | | |

REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSMER 70 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN

ACCESSION NUMBER: 1998:664773 CAPLUS

DOCUMENT NUMBER: 10:13971

TITLE: 10:13971

New synthesis of alloxazine derivatives

Krasnov, K. A.

CORPORATE SOURCE: State Chemical and Pharmaceutical

Academy, St. Petersburg State Chemical and Pharmaceutical

Academy, St. Petersburg, 1973/6, Russia

Russian Journal of Organic Chemical (1998), 34(1), 115-119

CODIN. RIDGE(1 ISSN: 1070-4280

PUBLISHER: MAIK Nauke/Interperiodica Publishing

JOURNAL

AMIK Nauke/Interperiodica Publishing

JOURNAL

AMIK

215866-02-7 CAPLUS Benzo[g]pteridine-7-carboxylic acid, 1,2,3,4-tetrahydro-1,3-dimethyl-2,4-dloxo-, ethyl ester, 5-oxide (9CI) (CA INDEX NAME) 215866-02-7

Title compds. I [R = H, OH, NH2; R1 = R2 = H; or R1R2 = :NR3; R3 = H, CO2R6, COR6, CON(R6)2, CH3OR7, CH3SR7; R4 = H, slkyl, alkyl-O, thicheterocycyl(n)1, (CH3CR1)Ahr, (CH:CR1)Ahr, CH3AR; R5 = alk(en/yn)yl, cycloalk(en)yl, heterocycyl(en)yl, aryl, heteroaryl, tweed systems, etc.; R6 = H, lower alkyl; R7 = H, lower alkyl, aralkyl, lower acyl, aroyl, heterocroyl; R8 = H, lower alkyl, R7 = H, lower alkyl, aralkyl, lower acyl, aroyl, heterocroyl; R8 = R, lower alkyl, R9 = H, R1002C, R100, H0, cyano, R10CO, OHC, lower alkyl, ON, Y1'Y2'N, R10 = alkyl, aralkyl, artlakyl, tyl, aralkyl; Y1', Y2' = H, alkyl, aryl, aralkyl; on one of Y1 and Y3 = acyl or aroyl and the other is as given; Ar = aryl or heterocryl; n = 0-21 and their pharmaceutically acceptable salts, prodrugs, N-oxides, hydrates, and solvates, are useful as Factor Xa inhibitors. For example, 4 (pyridin-3-yl)benzoic acid was amidated with tert-Bu 3-aminopropionate-HCl via the acid chloride, and the resulting p-acylamine ester underwent a sequence of (1) u-alkylation with S-iodo-2-[(2-msthoxyethoxy)methoxylbensyl bromide, (2) acidic deprotection of the MM group, and conversion to the Me ester, (3) Pd-catalyzed cyanation of the iodide, and (4) Pinner reaction and ammonolysis of the nitrila, to give title compound II. Three example compds, showed Ki values of the MM in a Factor Xa mssss, 45 mN to 1.72 µM in a trypsin action, 3-10 ph in a trypsin (aminomethyl)- or (aminomethyl)- or ((aminomethyl)-ph) ph phylpropyl amides as Factor Xa inhibitors) 219673-02-6 CAPLUS
Bentenepropanoic acid, 3-(aminominomethyl)- or (1-(4c innosminomethyl) aminolethyl)-, methyl ester (SCI) (CA index NAME)

21967-07-6 CAPLUS Senzenepropanoic acid, 3-(aminoiminomethyl)- a-[1-[(6-quinoxalinylcarbonyl)amino]ethyl]-, methyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 71 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER: 1998:608600 CAPLUS
DOCUMENT NUMBER: 129:210740
TITLE: Heteroary1-hexanoic acid amide d

129:230740
Heteroaryl-bexanoic acid amide derivatives, their preparation and their use as selective inhibitors of MIP-1 a binding to its CCR1 receptor Brown, Matthew Frank; Kath, John Charles; Poss, Christopher Stanley Pfizer Inc., USA PCT Int. Appl., 106 pp. CODEN: PIXXD2
Patent

INVENTOR (S) :

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PA | TENT | NO. | | | KIN | D | DATE | | А | PPL | ICAT | ION | NO. | | 1 | DATE | |
|---------|--------------------------------------|-----|------|-----|-----|-----|-------|------|-----|-----|---------------|------|-----|-----|-----|-------|-----|
| | | | | | | | | | | | | | | | | | |
| WO | 9838 | | | | | | | | | | | | | | | | |
| | W: | | | | | | | | BG, | | | | | | | | |
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| | | | | | Yυ, | | | | | | | | | | | | |
| | RW: | | | | | | | | | | | | | | | | |
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| | | GΑ, | GN, | ML, | MR, | NE, | , SN, | TD, | TG | | | | | | | | |
| CA | 2282 | 834 | | | AA | | 1998 | 0903 | C | A 1 | 998- | 2282 | 834 | | 1 | 9980 | 205 |
| CA | 2282 | 834 | | | C | | 2004 | 1005 | | | | | | | | | |
| AU | 9861 | 354 | | | A1 | | 1998 | 0918 | A | U 1 | 998- | 6135 | 4 | | 1 | 19980 | 205 |
| AU | 2282
2282
9861
7456
9664 | 67 | | | B2 | | 2002 | 0328 | | | | | | | | | |
| EP | 9664 | 43 | | | A1 | | 1999 | 1229 | Е | P 1 | 998- | 9060 | 13 | | 1 | 19980 | 205 |
| | R: | | | | | | | FR, | GB, | GR, | IT, | LI, | LU, | NL, | SE, | PT, | IE, |
| | | | | | FI, | | | | | | | | | | | | |
| TR | 9902 | 056 | | | T2 | | 2000 | 0121 | T | R 1 | 999- | 9902 | 056 | | 1 | 9980 | 205 |
| | 9607 | | | | | | | | | | | | | | | | |
| | 2000 | | | | | | | | | | | | | | | | |
| JP | 3771 | 591 | | | B2 | | 2006 | 0426 | _ | | | | | | | | |
| IL | 3771
1311
9801
1056 | 63 | | | A1 | | 2005 | 0619 | 1 | L 1 | 998- | 1311 | 63 | | 1 | 9980 | 205 |
| ZA | 9801 | 603 | | | | | 1999 | 0931 | 2 | A 1 | 998- | 1602 | | | - 1 | 9980 | 726 |
| AP | 1056 | | | | | | 2002 | 0405 | A | P 1 | 998- | 1200 | | | 2 | 9980 | 226 |
| | W: | | | | | | | | | | | | | | | | |
| | 1036 | | | | | | | | | | | | | | | | |
| | 9904 | | | | | | | | | 0 1 | 999- | 4101 | | | | 9990 | 825 |
| NO | 3138 | 77 | | | B1 | | 2002 | 1216 | | | | | | | | | |
| US | 6403 | 587 | | | 81 | | 2002 | 0611 | U. | 9 2 | 000- | 3802 | 69 | | - | 0000 | 518 |
| | 2002 | | | | Al | | 2002 | 1226 | U, | 3 2 | 002- | 1541 | 45 | | . 3 | 0020 | 522 |
| PRIORIT | Y APP | ښN. | INFO | . : | | | | | U | 9 1 | 997-
998-1 | 3916 | 9 P | | , 1 | 9970 | 226 |
| | | | | | | | | | W | . 1 | 338- | U215 | 8.6 | | | 9980 | 205 |
| | | | | | | | | | Մ | 5 2 | 000- | 3802 | 69 | | u a | 0000 | 218 |
| | | | | | | | | | | | | | | | | | |

OTHER SOURCE(S): MARPAT 129:230740

I [R1 = optionally substituted (C2-C9)heteroary1; R2 = optionally
substituted pheny1-(CH2)m-, naphthy1-(CH2)m-, (C3-C10)cycloalky1-(CH2)m-,
(C1-C6)alky1 or (C2-C9)heteroary1-(CH2)m-; m = integer from zero to four;
R3 = R, optionally substituted (C1-C10)alky1, (C3-C10)cycloalky1-(CH2)n-;
(C2-C9)heterocycloalky1-(CH2)n-, (C2-C9)heteroary1-(CH2)n-, ary1-(CH2)n-;
n = integer from zero to six; R1 and the carbon to which it is attached
form an optionally substituted and/or fused five to seven membered
carbocyclic ring; R8 = H, (C1-C6)alky1, dydroxy, (C1-C6)alkxy1-,
hydroxy-(C1-C6)alky1, (C1-C6)alkoxyC0, (C3-C10)cycloalky1-(CH2)p-,
optionally substituted (C2-C9)heterocycloalky1-(CH2)p-, p = integer
from zero to four; R4 and R5 together with the nitrogen atom to which they
are attached form an optionally substituted (C2-C9)heterocycloalky1 group;
R5 = H, (C1-C6)alky1, mainol were prepared The present compds are potent
and selective inhibitors of MIP-1 α binding to its receptor CCR1, and
are thus useful to treat inflammation and other immune disorders. E.g.,
quinoxaline-2-carboxylic acid [1(8)-benzyl-4(R)-benzylcarbamoyl-7-fluoro1(8)-hydroxy-7-methyl-ctyllamide wesp prepared
212789-54-20-200.
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12789-54-30-200.
12

Absolute stareochemistry.

REFERENCE COUNT:

THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 72 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DICCUMENT NUMBER:
119:819402403 CAPLUS
129:81940 Mee
129

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PAT | ENT | NO. | | | KIN | D | DATE | | | APPL | ICAT | ION | NO. | | D | ATE | |
|-----|------------|-----|-----|-----|-----|-----|------|------|-----|------|------|------|-----|-----|-----|------|-----|
| | | | | | | - | | | | | | | | | - | | |
| WO | WO 9825883 | | | | A1 | | 1998 | 0618 | 1 | WO 1 | 997- | EP66 | 55 | | 1 | 9971 | 126 |
| | W: | AL, | Aυ, | BG, | BR. | BY, | CA, | CN. | CZ. | GE, | HU, | ID. | IL. | JP, | KR. | KZ, | LT. |
| | | LV, | MX. | NO, | NZ. | PL. | RO, | RU. | SG. | SI. | SK. | TR. | UA. | US. | AM. | AZ. | BY. |
| | | KO. | KZ. | MD. | RU. | TJ. | TM | | | | | | | | | | |
| | | | | | | | | | | | | | | | | | |

209174-23-2 CAPLUS 6-Quinoxalinecarboxamide, N-[[4-{[[[18]-3-amino-2-hydroxy-3-oxo-1-(phenylmethyl)propyl]amino]carbonyl]phenyl]methyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

3

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 73 OF 1e1 CAPUUS COPYRIGHT 3006 ACS on STN
ACCESSION NUMBER: 1993-46203 CAPUUS
COPYRIGHT 3006 ACS on STN
1993-46203 CAPUUS
TITLE: 129.14620 CAPUUS COPYRIGHT 3006 ACS on STN
AUTHOR (S): 129.14620 CAPUUS CAPUU

Initial concentration ine values obtained were magn and comparable for error and Levafix dyes (5.4 and 5.7-106 M-1, resp.) and significantly smaller for Reactive Orange 046 and Reactive Red 022 (0.23 and 0.4-106 M-1). A possible interpretation of these data, based on the different structures of the bonded dyes, is proposed.

106058-73-1
RLI ARG (Analytical reagent use); PEP (Physical, engineering or chemical process); ANST (Analytical study); PROC (Process); USES (Uses) (Levafix Brilliant Blue S-B; reactive dyes-human serum albumin binding by frontal anal. affinity chromatog.)

2-Anthracenesulfonic acid, 1-maino-4-{[[4-[[(2,3-dichloro-6-quinoxaliny])esrbony]]smino]mothyl]-3-mulfophenyl]methyl]mmino]-9,10-dihydro-9,10-dioxo-, disodium selt (9CI) (CA INDEX NAME)

| RW: | : AT, BE, | CH, DE, | DK. ES. FI. | FR, GB, GR, IE, IT, | LU, H | C, NL, PT, | SE |
|--------------|-----------|---------|-------------|---------------------|-------|------------|----|
| CA 2274 | 1464 | AA | 19980618 | CA 1997-2274464 | | 19971128 | |
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| EP 9445 | 582 | B1 | 20030702 | | | | |
| R: | AT. BE. | CH. DB. | DK. ES. FR. | GB, GR, IT, LI, LU, | NL. S | E. PT. IE. | |
| | SI. PI. | | | | | | |
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| BG 6338 | 12 | B1 | 20011231 | BG 1999-103465 | | 19990611 | |
| PRIORITY API | |). : | | DE 1996-19651316 | A | 19961211 | |
| | | | | WO 1997-EP6655 | W | 19971128 | |
| | | | | | | | |

RETY APPEN. INFO::

RE SOURCE [8]:

WARPAT 139:13964

RE SOURCE [8]:

WARPAT 139:13964

The invention concerns ketobensamides of formula RIX(R2)n-CSH1COHHER(R2)COCR8 [1.2] (MR5, CO2H, Chebell (Mesteriotycle) (Pateriotycle), R2 COHHER(R2)COCR8 [1.2] (MR5, CO2H, [substituted]-slky], -slkenyl, -slkynyl,
R5 = CO-slkyl, CO2h, CO-C10H7, SO2-slkyl, CO2h, calkoxy, ureids, slkoxy; R3 =
(substituted) slkyl; X = (substituted) (functionalized)chain from 0-10

stoms, or R2-substituted-CSH3; R4 = OM, (substituted)lakoxy,
(substituted)NH2, heterocyclic ringl, useful as calpsin inhibitors. The
invention further concerns their preparation The novel compds. are suitable
for combsting diseases. Thus, 3(S)-3-mino-2-hydroxy-4-phenylbutyric acid
Me ester was condensed with 2-phenylbenzoic acid to give [8]-1 [R1 * Ph; X
- null; n = 0; R3 = CH2Ph; R4 = OMe(II)]. In in vitro calpsin-inhibition
tests, II had KI of <10 µM.
209174-24-39

RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study), unclassified); SPN (Synthetic preparation); BIOL (Biological
study); reparation end use of ketobenzamides as calpsin inhibitors)
20174-24-33 CAPLUS OTHER SOURCE(S):

209174-24-3 CAPLUS
6-Quinoxalinearboxamide, N-[[4-[([18]-3-amino-2,3-dioxo-1-(phenylmethyl)propyl]amino|carbonyl|phenyl|methyl]- [9CI) (CA INDEX NAME)

Absolute stereochemistry.

209174-23-2P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) IT actant or reagent) (preparation and use of ketobenzamides as calpain inhibitors)

PAGE 1-A

PAGE 2-A

THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT REFERENCE COUNT:

3 ANSWER 74 OF 181 CESSION NUMBER: CAPLUS PLUS COPYRIGHT 2006 ACS on STN 1998:268491 CAPLUS

DOCUMENT NUMBER: TITLE:

1999:26491

128:30499

Bis (acridinecerboxamide) and bis (phenazinecerboxamide) as antitumor sgents

Denny, William Alexander; Gemage, Swarmaletha

Akuritays; Spicer, Julie Ann; Baguley, Bruce Charles;

Finley, Oraeme John

Kenova Led. UK, Denny, William Alexander; Gamage,

Swarmalatha Akuritays; Spicer, Julie Ann; Baguley,

Bruce Charles; Finley, Graeme John

PCT Int. Appl., 100 pp.

CODEN: PIXXO2

Patent

Rollish

PATENT ASSIGNEE(S):

SOURCE:

DOCUMENT TYPE: PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. APPLICATION NO. DATE KIND DATE MO 9817650 A1 19980430 MO 1997-GB2886 19971017 M: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DR,

| | | DK, | EE, | ES, | FI. | GB, | GE, | GH, | HU, | ID | , IL. | IS, | JP. | KE, | KO, | K₽, | KR, |
|---------|-------|------|-----------|-----|-------|------|------|------|------|-----|--------|------|--------|------|-----|--------|-----|
| | | KZ. | IC. | LK, | LR, | LS. | LT. | LU. | LV. | Ю | , MO, | MX, | MN, | MNF, | MX. | NO. | NZ. |
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| | | us, | υz, | VN, | YU, | ZW | | | | | | | | | | | |
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| ZA | 97093 | 331 | | | λ | | 1998 | 0521 | : | ZA | 1997- | 9331 | | | 1 | 9971 | 017 |
| ZA | 9709 | 328 | | | Α | | 1998 | 0706 | : | ZA | 1997- | 9328 | | | 1 | 9971 | 017 |
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| EP | 9342 | 76 | | | | | | | | | | | | | | | |
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FI | CH, | DE, | DK, | ES, | FR, | GB, | GR | , іт, | LI, | LU, | NL, | SE, | MC, | PT. |
| GB. | 23340 | | •1 | | A1 | | 1000 | 0811 | | 20 | 1999- | 1100 | | | | 0071 | 017 |
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| ON. | 12404 | 130 | | | â | | | 0105 | | | 1997- | | | | | 9971 | |
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| | 33509 | | | | Ä | | | 0929 | | NZ. | 1997- | 3350 | 44 | | | 9971 | 117 |
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| | 93427 | | | | Ī | | | 0131 | - 1 | PT | 1997- | 094 | 56 | | 1 | 9971 | 017 |
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| | 61143 | | | | Ä | | | 0905 | | | 1999- | | | | | | |
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| IORIT | | | | | | | | | | | 1996- | | | | | | |
| | | | | | | | | | | | 1997-0 | | | | | 971 | |
| THER SO | URCE | (R): | | | CASI | REAC | F 12 | 8:30 | 1499 | . M | ARPAT | 128 | : 3084 | 199 | | | |
| | | | | | | | | | | | | | | | | | |

Compds. I [R1-R4 = H, C1-4 alkyl, OH, etc.; or R1 and R2 together form a methylenedioxy group; R5, R6 = H, C1-4 alkyl; X = CH, N; Z = (CH2)n, (CH2)nOR(CH2)n, (CH2)nOR(CH2)n, (CH2)nOR(CH2)n, RCH2)nNPT(CH2)n, (CH2)nNPT(CH2)n, (CH2)nNPT(CH2)n, R7 = H, C1-4 alkyl; m, n = 1-4; with the exception of compds. wherein each K is N, each of R1-R6 is H, the carboxamide moiety is attached to position 1 of each phenazine ring and Z is (CH2)2NNH(CH2)2, CH2)3NH(CH2)3, (CH2)3NH(CH2)3, (CH2)3NH(CH2)3, (CH2)3NH(CH2)3, CH2)3NH(CH2)3, or (CH2)2NH(CH2)3, coreptable acid addition salt or N-oxide thereof; have activity as an antitumor and antibacterial agent. Thus, bis[15-methylacridine-4-carboxamido)propyl]sethylamine was prepared and showed an IC50 value of 11 nM on a wild-type human leukemia line (Jurkat; JLc).

PAGE 2-A

PAGE 1-A

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE PORMAT

L13 ANSWER 76 OF 181 CAPUUS COPYRIGHT 2006 ACS on STN
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DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

REPERENCE COUNTY

NO. KIND DATE APPLICATION NO. DATE

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AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DB,
KE, ES, FI, GB, GE, GH, HU, ID, IL, IS, JP, KE, KG, KP, KR,
KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, NN, NM, MK, NO, NZ,
PL, FT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TH, TR, TT, UA, UG,
US, UZ, VN, YU, ZM, AN, AZ, BY, KG, KZ, KG, NU, TJ, TM PATENT NO. WD 9813368

IT 20531-48-8P
RL: SPN (Shrhetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(bis(acridinecarboxamide) and bis(phenasinecarboxamide) as antitumor and antibacterial agents)
RN 206531-48-8 CAPLUS
CN 2-Phenasinecarboxamide, N,N'-[(methylimino)di-3,1-propanediyl]bis-, monohydrochloride (9CI) (CA INDEX NAME)

● RC1

REFERENCE COUNT: THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE PORMAT

L13 ANSWER 75 OF 181 CAPLUS COPYRIGHT 2006 ACE on STN
ACCESSION NUMBER:
1998:266807 CAPLUS
120:295812
An improved computational approach to the
determination of thermodynamic and spectral
complexation parameters from overlapping bands.
Applications to the case of the Cibacron dimer and to
the multiple HSAV-Levafix association
AUTHOR(8):
RODERORATE SOURCE:
CORPORATE SOUR

CORPORATE SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE:

Mores (8):

Ambrosetti, Roberto; Ricci, Domenico; Bianchini, Roberto
PORATE SOURCE:

CMR. Instituto di Chimica Quantistica ed Energetica
Molecolare, Pisa, I-56126, Italy
RCE:

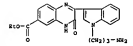
Gazzetta Chimica Italiana (1997), 127(10), 567-575
CODEN: OCITA9; ISSN: 0016-5603
Societa Chimica Italiana
DURINT TYPE:

JOURNEL JOURNEL
An algorithm for the simultaneous evaluation of the thermodn. parameters
related to multiple equilibrium and of the spectra of dya complexes is
described. The algorithm can accept as input data any stochiometry for
complex species and relies on the simultaneous fitting of large sets of
data obtained at different concns., temps., and wavelengths. Data from
different measuring techniques, such as UV-visible absorption or CD, may
be included in a single fit. Datails on an easily modifiable, yet
computationally efficient implementation of the algorithm on a standard PC are
given. Results are presented for the dimer aggregation of the dya
Cibacron Blue F3GA and for the association of human serum albumin
RL: PRP (Properties)
(algorithm for determination of spectral and thermodn. parameters of)
206058-73-3 CAPUUS
2-Anthracenesulfonic acid, 1-amino-4-[[[4-[[[(2,3-dichloro-6quinoxalinyl]carbonyl]amino]methyl]-3-sulfophenyl]methyl]amino]-9,10dihydro-9,10-dioxo-, disodium salt (9CI) (CA INDEX NAME)

RM: GH, KE, LS, MM, SD, SZ, UG, ZW, AT, BB, CH, DE, DK, ES, FT, FR,
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IE, SI, LT, V, FT, RG
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KTTY APPLN, INFO: SE 1996-3505 A 19960925 NZ 1997-334531 US 1997-981266 US 2001-865231 SE 1996-3505 SE 1997-2747 WO 1997-SE1582 US 1997-981266 NZ 334531 US 6271231 US 2001025043 PRIORITY APPLN. INFO.: 19971218 20010525 A 19960925 A 19970718 W 19970919 A3 19971218 OTHER SOURCE(S): MARPAT 128:257445

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CRN 205376-67-6 CMP C22 H22 N4 O3



CRN 64-19-7 CMF C2 H4 O2

REPERENCE COUNT:

THERE ARE 12 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 77 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1998:126254 CAPLUS
DOCUMENT NUMBER: 128:204878
TITLE: Preparation of pyrazinobenzothia

INVENTOR (S) :

128:204878

Preparation of pyrazinobenzothiazine derivatives and analogs for the treatment of inflammation and autoimmune diseases

Kaneko, Toshihiko; Clark, Richard; Ohi, Norihito;
Ozaki, Pumihiro; Kawahara, Teteuya; Kamada, Atsushi;
Okano, Kazuo; Yokohama, Hiromitau; Muramoto, Kenzo;
Arai, Tohru; Ohkuro, Masayoshi; Takenaka, Osamu; Arai, Tohru; Ohkuro, Mas Sonoda, Jiro Bisai Co., Ltd., Japan PCT Int. Appl., 1344 pp. CODEN: PIXXD2 Patent Japanese

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: FAMILY ACC. NUM. COUNT: 1

WO 9736907

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| | PA | TE | T : | NO. | | | KIN | D | DATE | | | AP | PLIC | AT | ION | NO. | | | DATE | | |
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| 10 | RIT | YA | PP | LN. | INFO |).: | | | | | | JΡ | 199 | 6-2 | 210 | 144 | | A | 19960 | 809 | |
| | | | | | | | | | | | | WO | 199 | 7- | JP2 | 787 | | w | 19970 | 808 | |
| | | | | | | | | | | | | US | 199 | 9-: | 230 | 52 | | A3 | 19990 | 405 | |
| HE | R S | OUR | CE | (S): | : | | MAR | PAT | 126: | 2048 | 78 | | | | | | | | | | |
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| | | | DK, | EE, | ES, | PI, | GB, | GE, | HU, | IL, | IS. | , J₽, | KE, | KG, | KP, | KR. | KZ, | LC, |
| | | | LK, | LR, | LS, | LT, | LU, | LV, | MD, | MG, | MK | , MN, | MW, | MX, | NO, | NZ, | PL, | PT, |
| | | | RO, | RU, | SD, | SE, | 5G, | SI, | 5Κ, | TJ, | TM | , TR, | TT, | UA, | VG, | UZ, | VN, | YU, |
| | | | AM, | AZ, | BY, | KG, | KZ, | MD, | RU, | TJ, | TM | | | | | | | |
| | | RW: | GH, | KE, | LB, | MW, | SD, | SZ. | UG, | AT, | BE | , СН, | DE, | DK, | ES, | FI. | FR, | GB, |
| | | | GR, | IE, | IT, | LU, | MC, | NL, | PT, | SE, | BF | BJ, | CF, | co, | CI, | CM, | GA, | GN, |
| | | | ML, | | | SN, | | | | | | | | | | | | |
| | US | 5736 | 543 | | | A | | 1998 | 0407 | - 1 | US : | 1996- | 6243 | 3 5 | | | 19960 | 403 |
| | CA | 2249 | 654 | | | AA | | 1997 | 1009 | | CA : | 1997- | 2249 | 654 | | | 19970 | 328 |
| | | | | | | | | | | | AU : | 1997- | 2427 | 4 | | | 9970 | 328 |
| | AU | 7082 | 13 | | | B2 | | 1999 | 0729 | | | | | | | | | |
| | EP | 8913 | 65 | | | A1 | | 1999 | 0120 | - 1 | BP : | 1997- | 9199 | 65 | | - 1 | 19970 | 328 |
| | EP | 8913 | 65 | | | B1 | | 2002 | 1016 | | | | | | | | | |
| | | R: | AT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, | GR | IT, | LI, | LU, | NL, | SE, | MC. | PT. |
| | | | IE, | PI | | | | | | | | | | | | | | |
| | JÞ | 2000 | 5036 | 64 | | T2 | | 2000 | 0328 | | JP : | 1997- | 5354 | 86 | | 1 | 19970 | 328 |
| | J₽ | 3170 | 294 | | | B2 | | 2001 | 0528 | | | | | | | | | |
| | AT | 2262 | 06 | | | | | 2002 | 1115 | - 1 | AT : | 1997- | 9199 | 65 | | 1 | 9970 | 328 |
| | 88 | 2185 | 932 | | | T3 | | 2003 | 0501 | 1 | ES : | 1997- | 9199 | 65 | | 1 | 19970 | 328 |
| | US | 5962 | 447 | | | A | | 1999 | 1005 | 1 | US : | 1998- | 1988 | 3 | | 1 | 19980 | 206 |
| | ΑU | 9944 | 543 | | | A1 | | 1999 | 1021 | - 1 | AU : | 1999- | 4454 | 3 | | 1 | 9990 | 817 |
| | ΑU | 7219 | 36 | | | 82 | | 2000 | 0720 | | | | | | | | | |
| RIC | RIT | Y APP | LN. | INFO | . : | | | | | 1 | US : | 1996- | 6243 | 35 | 1 | A 1 | 9960 | 403 |
| | | | | | | | | | | 1 | AU : | 1997- | 2427 | 4 | - 1 | A3 1 | 9970 | 326 |
| | | | | | | | | | | 1 | NO : | 1997- | US51 | 84 | 1 | ¥ 1 | 9970 | 328 |
| THE | R SC | URCE | (8): | | | MARP | AT | 127: | 3189 | 71 | | | | | | | | |

19921009 WO 1997-USE184

Compds. of formula [I; Rl, R2 = independently selected from the group consisting of H and R40 such that at least one of R1 and R2 is R40; R4 = a member selected from the group consisting of H, alkyl and halo-substituted alkyl; or R1 and R2 together form a single divalent moiety selected from the group consisting of O-R5-0, S-R5-0, O-R5-8, N:CR6CR7:N, O-CR8:N, N:CR80; Wherein R5 = a member selected from the group consisting of C(R9)2, CR9)2c(R9)2, CR9:CR9; R9 = H, halo, C1-6 alkyl, C1-6 haloslkyl; R3, R6, R7, R8 = a member selected from the group consisting of R, C1-6 alkyl, and C1-6 haloslkyl] are disclosed for use in enhancing synaptic responses mediated by a-anino-1-hydroxy-5-methyl-4-isoxazolepropionic (AMPA) receptors. The compds. are effective in the treatment of subjects suffering from impaired nervous or intellectual functioning due to deficiencies in the number or strength of excitatory synapses or in the number of AMPA receptors. The compds. can also be used for the treatment of non-impaired subjects for enhancing performance in sensory-monter and cognitive tasks which depend on brain networks utilizing AMPA receptors, for improving the performance of subjects with memory encoding. Thus, 3,4-methylenedioxysalicylic acid was condensed with 4-aminobutyraldehyde di-Rt acetal using N,N-carbonyldimidazole in CH3C12 at room temperature overnight to quant, give N-(4,4-disthoxybutyl)-1,4-methylenedioxysalicylanide, which was dissolved in CRC13 and allowed to stand in the presence of camphorsulfonic acid overnight to give I (R1R2 = OCH2CH2O, R3 = H, n = 2). The latter compound and I (R1R2 = OCH2CH2O, R3 = H,

$$\begin{array}{c|c}
R & R^1 \\
R^2 & R^3
\end{array}$$

The title compds. I [R1 to R3 are the same or different and each represents hydrogen, optionally substituted lower slkyl, optionally substituted cyclealkyl, etc., provided that when R1 to R3 are all optionally substituted lower alkyl groups, they do not simultaneously represent Me groups; R represents hydrogen, lower alkyl, etc.; S represents N. C, etc.; Z represents O, S, SO, SO, etc.; and the ring G represents an optionally substituted heteroaryl ring having at least on nitrogen atom) are prepared I are useful in the treatent and pravention of inflammatory immunol. diseases, autoimmune diseases, rhoundatism, collegen disease, sathums, nephritis, ischemic reflow disorders, psoriasis, atopic dermatitis or rejection reactions following organ transplantation. The compound (sym): [3-10H-pyrazino[3-3-b][1, 4]benzothisain-8-ylmethyl)-3-azabicyclo[3.3.1]nons-9-yllacetic acid [II] at 10 mg/kg orally gave 651 inhibition of carragenin-induced inflammation in rate. II in vitro showed IC50 of 2.3 µM against the expression of ICAM-1.

showed ICSO of 2.3 µM against the expression of ICAM-1.
201651-75-6P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological atudy); PREP (Preparation); USES (Uses)
(preparation of pyrazinobenzothiazine derive. and analogs for treatment of inflammation and autoimmune diseases)
203651-75-6 CAPLUS
Pyrazino[2,3-b]quinoxaline-7-carboxylic acid, 1.4-dihydro-, ethyl ester
(9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 46 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 46

L13 ANSWER 78 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1997:679093 CAPLUS DOCUMENT NUMBER: 127:318971

TITLE: Preparation of benzoxazinea for enhancing synaptic

response
Rogers, Gary A.; Lynch, Gary S.
Regents of the University of California, USA; Cortex
Pharmaceuticals, Inc
PCT Int. Appl., 34 pp.
CODEN: PIXXD2
Patent
English
1 INVENTOR(S): PATENT ASSIGNEE(S):

SOURCE:

DOCUMENT TYPE:

PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

n = 2) in vivo showed the threshold dose of 1 and 0.1 mg/kg, resp., for enhancing memory in rate in a learning paradigm that depends on a performance in an 8-arm radical maze described by Staubli et al. (PMAS, 1994), and at 0.1 and 0.03 md in vitro, resp. increased the amplitude of the field excitory poet-synaptic potential (EPSP) to a value 25% above the base line in slices of rat hippocampus.
197584-95-65 197589-97-79
RL: RCT (Resectant); SPM (Synthetic preparation); PREP (Preparation); RACT (Reactant or resgent)
(preparation of benzoxazines with affinity to AMPA receptors for enhancing synaptic response)
197584-96-6 CAPLUS
6-Quinozalinecarboxamide, N-(4,4-diethoxybutyl)-7-hydroxy- (9CI) (CA INDEX NAME)

197584-97-7 CAPLUS 6-Quinoxalinecerboxamide, N-(5,5-diethoxypentyl)-7-hydroxy- (9CI) (CA INDEX NAME)

L13 ANSWER 79 OP 181 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:
DOCUMENT NUMBER:
1155:259566

Electrochemical study of C60-containing dimine and polyazine ligands: towards a fullerone-besed photoactive molecular device

AUTHOR(S):

AUTHOR(S):

AUTHOR(S):

Paradisi, Carmen, Reffis, Sergio; Prato, maurizio
Dep. Chem., Univ. Bologna, Bologna, 40126, Italy
Proceedings - Slectrochemical Society (1996),
96-10(Racent Advances in the Chemietry and Physica of Pullerones and Related Materials, Vol. 3), 157-164

CODEN: PRSODO; ISSN: 0161-6374

Blectrochemical Bociety
Journal
LANGUAGE:
English

ANGUAGE Tree:

Anguage The complete electroches. characterization of the fulleropyrrolidines containing the 2,2'-bipyridine (bpy) and the 2,3-bis(2-pyridyl)quinoxaline (dpq) fragment, resp., is reported. These ligands represent important building blocks for the assembling of supremol. (polynuclear) metal complexes in which a photoinduced intramol. charge exparation may in principle take place. The comperison of the cyclic voltammograms of these species with those of suitable model mole. has allowed the localization of the eight reversible reduction processes observed for both species, a fundamental polynuclear complexes.

IT 182219-47-2 LANGUAGE: English

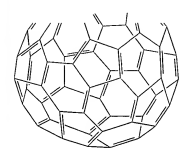
RL: PEP (Physical, engineering or chemical process); PRP (Properties); RCT (Reactant); PROC (Process); RACT (Reactant or reagent) (electrochem. reduction in THF containing tetrabutylammonium hexafluorophosphate: towards fullerene-based photoactive mol. device) 18219-47-2 CAPLUS 6-Quinoxalinecarboxylic acid. 3,3-di-3-pyridinyl-, 13-[4-[1',5'-dihydro-1'-methyl-2'H-[5,6]fullereno-C60-lh-[1,9-c]pyrrol-2-yl)phenyl]-13-oxo-3,6,9,12-tetraoxatridec-lyl ester (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

PAGE 1-A

PAGE 1-B

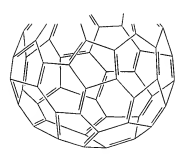


PAGE 3-A

PAGE 2-A



182219-47-2D, transition metal complexes
RL: PEP (Physical, engineering or chemical process); PRP (Properties);
PROC (Process)
(photoinduced intramol. charge separation in)
182219-47-2 CAPLUS
6-Quinoxalinecarboxylic scid, 2,3-di-2-pyridinyl-, 13-[4-[1',5'-dihydro-1'-methyl-2'H-[5,6](fullereno-C60-Ih-[1,9-c]pyrrol-2-yl]phenyl]-13-oxo3,6,9,12-tetraoxatridec-1-yl ester (9CI) (CA INDEX NAME)



PAGE 2-A

PAGE 3-A

LI3 ANSWER 80 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
1996:\$42076 CAPLUS
DOCUMENT NUMBER:
125:204109
Silectrochemical and SSR spectroscopic study of
2,7-disubstituted phenarines
AUTHOR(8):
AUTHOR(8):
Fac. Pharmacctuical Sci., Kobe-Gakuin Univ., Kobe,
651-21, Japan
COURCE:
Chemical & Pharmaccutical Bulletin (1996), 44(8),
1448-1453
COODN: CPSTAL; ISSN: 0009-2161
PUBLISHER:
Pharmaccutical Society of Japan
DOCUMENT TYPE;
Journal
LANGUAGE:
Beglish
AB Cyclic voltammetry (CV) for various 2,7-disubstituted phenazines (1 mM)
was carried out in MeCN containing CTSCOOK (14 and 24) and NaClO4 (0.1M) as a
supporting electrolyte under N. Phenazines showed 2 cathodic peaks (Epc1
and Epc2) and these peaks had counterparts (Eps1 and Spc2, resp.). Plots
of the peak potentials against op were linear. The first cathodic
wave corresponds to the reduction of singly protonated phenazines followed by
proton transfer. The second cathodic wave corresponds to the reduction of the
cation radical of dihydrophenazines to produce dihydrophenazines followed by
product. ESR spectrometry of these compds. in MeCN and in MeCN containing 13
CF3CO2H was conducted and computer simulation of the spectra was carried

out. Splitting due to halogen or o-alkyl substituents was observed MO calcn. of anion radicals generated from the phenazines and cation radicals generated from doubly protonated phenazines did not give good agreement with the results of ESR spectrometry.

72848-45-4
RL: PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(electrochem. and ESR spectroscopic study of disubstituted phenazines)

72848-45-4 CAPLUS

2,7-Phenazinedicarboxylic acid, diethyl ester (9CI) (CA INDEX NAME)

L13 ANSWER 81 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1996:170757 CAPLUS
DOCUMENT NUMBER: 124:202310
TITLE: Preserved: 1

124:202310
Proparation of 1,2,3,4-tetrahydro-2,3-dioxoquinoxaline-6-sulfonemides as AMPA and kainate receptor antagonists
Rivo, Endre; Vizi, E. Szilveszter; Makara, Gabor; Reiter, Jozsef; Blasko, Gabor; Simig, Gyula; Gaal, Laszlo; Pekte, Marton
Egis Gyogyszergyar Rt., Hung.
PCT Int. Appl., 24 pp.
CODSN: PIXXD2
Patent

PATENT ASSIGNEE (S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---|------------------------|----------|-----------|---------------------|-----------------|
| | | | | | |
| | WO 9531443 | A1 | 19951123 | WO 1995-HU15 | 19950518 |
| | W: AM, AT, AU | , BB, BG | , BR, BY, | CA, CH, CN, CZ, DE, | DK, EE, ES, FI, |
| | GB, GB, HU | , IS, JP | , KE, KG, | KP, KR, KZ, LK, LR, | LT, LU, LV, MD, |
| | MG, MN, MW | , MX, NO | , NZ, PL, | PT, RO, RU, SD, SE, | SG, SI, SK, TJ, |
| | TT, UA | | | | |
| | RW: KE, MW, SE | , sz, ug | , AT, BE, | CH, DE, DK, ES, FR, | GB, GR, IE, IT, |
| | LU, MC, NI | , PT, SE | , BF, BJ, | CF, CG, CI, CM, GA, | GN, ML, MR, NE, |
| | SN, TD, TG | | | | |
| | HU 71334 | A2 | 19951128 | HU 1994-1522 | 19940518 |
| | HU 217837 | В | 20000428 | | |
| | CA 2190532 | AA | 19951123 | CA 1995-2190532 | 19950518 |
| | AU 9524162 | A1 | 19951205 | AU 1995-24162 | 19950518 |
| | EP 759910 | A1 | 19970305 | RP 1995-916107 | 19950518 |
| | R: AT, BE, CH | , DE, DK | , ES, PR, | GB, GR, IT, LI, NL, | SE |
| | JP 10504019 | T2 | 19980414 | JP 1995-529476 | 19950518 |
| | US 5912245 | A | 19990615 | US 1997-737273 | 19970214 |
| 1 | PRIORITY APPLN. INFO.: | | | HU 1994-1522 | A 19940518 |
| | | | | WO 1995-HU15 | W 19950518 |
| • | THER SOURCE(S): | MARPAT | 124:2023 | 10 | |
| • | 31 | | | | |
| | | | | | |

170467-25-19
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
170467-25-1 CAPLUS
6-Quinoxalinecarboxylic acid, 2-[[4-(butoxycarbonyl)phenyl]amino]-3-(1-methyl)-2-nitroethenyl)-, butyl ester (9CI) (CA INDEX NAME)

L13 ANSMER 83 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1995:716813 CAPLUS
DOCUMENT NUMBER: 123:112079
TITLE: Preparation of quinoxaline-2-carboxamides as antidiabetics
INVENTOR(S): Komateu, Makoto; Sato, Hideaki; Taira, Shinichi; Miyake, Masahiro; Magata, Kiyohiko; Yoshida, Hidehiro; Ueyama, Ateumori; Nishi, Takos
DOCUMENT ASSIGNEE(S): Octube Pharmaceutical Co. Ltd., Japan
PCT Int. Appl., 507
CODEN: PIXKD2
DOCUMENT TYPE:

Patent

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | • | | | | | | | | | | | | | | | | | |
|-----|-----|------|------|------|-------|-----|-----|-------|------|--------|-------|-------|-----|-----|-----|------|-----|----|
| 1 | PAT | TENT | NO. | | | KIN | D | DATE | | AP | PLICA | TION | NO. | | D | ATE | | |
| | | | | | | | | | | | | | | | - | | | |
| 1 | WO. | | | | | | | 1995 | 0406 | WO | 1994 | -JP15 | 59 | | 1 | 9940 | 922 | |
| | | W: | AU | , CA | CN, | KR, | us | | | | | | | | | | | |
| | | RW: | : AT | , BE | , сн, | DE, | DK. | , ES, | FR, | GB, GI | R, 16 | , IT, | LU, | MC, | NL, | PT, | SE | |
| - (| ZA. | 2150 | 345 | | | AA | | 1995 | 0406 | CA | 1994 | -2150 | 345 | | 1 | 9940 | 922 | |
| - 1 | AU | 9476 | 660 | | | A1 | | 1995 | 0418 | AU | 1994 | -7666 | 0 | | 1 | 9940 | 922 | |
| | ٩U | 6746 | 13 | | | 92 | | 1997 | 0102 | | | | | | | | | |
| 1 | 8P | 6708 | 331 | | | A1 | | 1995 | 0913 | EP | 1994 | -9270 | 85 | | 1 | 9940 | 922 | |
| | | R: | AT | . BE | CH, | DE. | DK. | ES. | FR. | GB, GI | R. IE | . IT. | LI. | LU. | MC. | NL. | PT. | SE |
| | 'n | 1114 | 834 | | | A | | 1996 | 0110 | CN | 1994 | -1907 | 19 | | 1 | 9940 | 922 | |
| į. | JP. | 0801 | 257 | 9 | | A2 | | 1996 | 0116 | JP | 1994 | -2593 | 09 | | 1 | 9940 | 928 | |
| | JP | 2759 | 257 | | | B2 | | 1998 | 0528 | | | | | | | | | |
| OR: | רדו | APE | LN. | INFO |).: | | | | | JP | 1993 | -2411 | 40 | | 1 | 9930 | 928 | |
| | | | | | | | | | | JP | 1994 | -1146 | 39 | | 1 | 9940 | 128 | |
| | | | | | | | | | | | | -JP15 | | | | | | |
| ER | 80 | URCE | (8) | : | | MAR | PAT | 123: | 1120 | | | | | | • | | | |
| | | | | | | | | | | | | | | | | | | |

OTHER SOURCE(S):

AB Title compds. (I; R = NR3R4; R1 = H, halo, alkyl, alkoxy, etc.; R2 = H,

AB

Title compds. [I; R1 = H or NO2; R2,R3 = H, (un)substituted alk(en)yl;
NR3R3 = heterocyclyl] were prepared Thus, I (R1 = NO2, NR3R3 = piperidino)
had Ki of 6.3x10-7 and 2.0x10-6M for inhibition of AMPA and Keinate
binding at rat brain membrane preparation in vitro.
174532-44-89
RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); SPN (Synthatic preparation); TNU (Theraputic use);
BIOL (Biological study); PREP (Preparation); USES (User)
(preparation of 1,2,3,4-tetrahydro-2,3-dloxoquinoxaline-6-sulfonamides as
AMPA and kainate receptor antagonists)
174536-64-8 CAPLUS
(Olycine, N-(1,2,3,4-tetrahydro-2,3-dloxo-6-quinoxalinyl)sulfonyll-,
phenylmethyl ester (9CI) (CA INDEX NAME)

M. Ross. Gos. Pedagog. Univ., Russia Zhurnal Organicheskoi Khimii (1994), 30(9), 1434-5 CODEN: ZORKAE; ISSN: 0514-7492 CORPORATE SOURCE:

SOURCE:

PUBLISHER: DOCUMENT TYPE: LANGUAGE: GI

Polynitro-3-thiolene 1,1-dioxide I reacted with para-substituted anilines to give quinoxalines (II; R = Me, Cl, Br, COOBu).

(halo)alkyl, alkoxy, etc.; R3,R4 = H, alkyl, alkanoyl, alkoxycarbonyl, substituted CH2Ph, heterocyclylalk(en)yl, etc.; m = 0 or 1; n = 0; r = 1 or 2] were prepared Thus, benzofuroxan was cyclocondensed with McCOCH2CO2Et and the product converted in 2 steps to I (R2 Me, m = 1, n = r = 0) (II; R = 0Et) which was amidated by 3-aminomethylbenzofuran to give II (3-benzofurylaminomethyl). II (R = NRCH2CHCRSMe, R5 = 2-benzofuryl) gave 2-deoxyglucose uptake of rat striated muscle L6 cells 249t of controls at 10-6mol (sic).
165735-35-3P
RL: BAC (Bological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); TRU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of quinoxaline-2-carboxamides as antidiabetics)
165735-35-3 CAPLUS
2,6-Quinoxalinedicarboxamide, N2-[3-(2-benzofuranyl)-2-butenyl]-N6,N6-diethyl-3-methyl-, 4-oxide (9CI) (CA INDEX NAME)

165736-35-67 165736-36-77 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation of quinoxaline-2-carboxamides as antidiabetics) 15736-35-6 CAPLUS 2-Quinoxalinecarboxylic acid, 6-((diethylamino)carbonyl)-3-methyl-, ethylester, 1,4-dioxide (DCI) (CA INDEX NAMS)

165736-35-7 CAPLUS
2-Quinoxalinecarboxylic acid, 6-[(diethylamino)carbonyl]-3-methyl-, ethylester, 4-oxide (9C1) (CA INDEX NAME)

L13 ANSMER 84 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1995:518899 CAPLUS
DECUMENT NUMBER: 133:6589 CAPLUS
13

PUBLISHER: DOCUMENT TYPE: LANGUAGE: GI

AB Hypoxic cells, which are a common feature of solid tumors, but not normal tissues, are resistant to both anticancer drugs and radiation therapy. Thus the identification of drugs with selective toxicity toward hypoxic cells is an important objective in anticancer chemotherapy. The benzotriazine di-N-oxide (SR 4233, Tirapazamine) has been shown to be an efficient and selective cytotoxin for hypoxic cells. Since the bioreductive activation of Tirapazamine is thought to be due to the presence of the 1,4-di-N-oxides with a range of electron-donating and -withdrawing substituents in the 6- and/or 7- postions has been synthesized and evaluated for toxicity to hypoxic cells. Electrochem. studies of the quinoxaline di-N-oxides and Tirapazamine showed that as the electron-withdrawing nature of the 6(7)-substituent increases, the reduction potential becomes more pos. and the compound is more readily reduced. Apart from the unsubstituted derivative and the 6,7-di-Me derivative I, the quinoxaline di-N-oxides have reduction potentials significantly more pos. than

The title compds. [I; A, B = cyano, NO2, halo, (un) substituted alkyl, alkenyl, alkynyl, aryl, alkyloxycarbonyl, aryloxycarbonyl, alkyloxycarbonyl, cyano, NO2; K, Y = H, (un) substituted alkyl, aryl, aryl, aryl, arylaxycarbonyl, cyano, NO2; K, Y = H, (un) substituted alkyl, aryl, aryl, arylaxycarbonyl, aryloxycarbonyl, alkylaminocarbonyl, or arylaminocarbonyll, having excellent electron-transport abhilty, are prepared An electrophtog, photoreceptor comprises a photosensitive layer containing 21 above compds. I as a charge-transport material, formed on a conductive support. Thus, 9.0 g ninhydrin and 7.6 g 3,4-diaminobenzoic acid was dissolved in SEOH and refluxed for 3 h to give a mixture of indenoquinoxalinone derivative (II; Z = O, R = CO2H, R1 = H) and regioisomer II (Z = O, R = H, R1 = CO2H) in 36t yield which (7.0 g) was exterified with BOOH in the presence of concentrated H2SO4 in refluxing toluene with removal of H2O through a ... Stark

concentrated BiSG4 in refluxing toluene with removal of H2O through a
-Stark
apparatus to give 11.9% Bu ester II (Z = 0, R = CO2Bu, Rl = H) and 70%
regioisomer II (Z = 0, R = H, Rl = CO2Bu). The latter regioisomer (0.4 g)
was refluxed with malonomitrile in the presence of piperidine in MeOH with
stirring for 14 h to give 97% title compound II (Z = C(M)2, R = H, Rl =
CO2Bu] (III). An electrophotog, photoreceptor with a charge-transport
layer containing III coated on an Al substrate was charged by a corona
discharge at +60 kV, left for 10 s in dark, and exposed with a 10-1x
halogen lamp to show maximum electrification potential (Ymax) of 420 V,
half-reduction exposure dose (EL/2) 15.0 l.s. and residual potential 110 V.
161290-68-69 161290-91-1P 161390-92-2P
RL: DEV (Device component use); SPN (Synthetic preparation); PREP
(Preparation); USES (Uses)
(preparation of indenoquinoxaline derivs. as charge-transport materials for
electrophotog, photoreceptors)
161290-68-6 CADLUS
11N-Indeno[1,2-b]quinoxaline-7-carboxylic acid, 11-(dicyanomethylene)-,
butyl ester (9CI) (CA INDEX NAME)

Tirapazamine (Epc -0.90 V). The most potent cytotoxins to cells in culture were the 6,7-dichloro and 6,7-difluoro derivs. II and III, which were 30-fold more potent than Tirapazamine. The 6(7)-fluoro and 6(7)-chloro compds. IV and V, showed the greatest hypoxia selectivity. Four of the compds. IV and V, showed the greatest hypoxia selectivity. Four of the compds. IV, VI, III and II, killed the inner cells of multicellular tumor spheroids in vitro. In vivo Balb/c mice tolerated a dose of these four compds. twice the eige of that of Tirapazamine. This study demonstrates that quinoxaline 1,4-di-N-oxides could provide useful hypoxia-selective therapeutic agents.
163777-45-59
RL: RAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PRPP (Properties); SFN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
(hypoxia-selective agents derived from quinoxaline di-N-oxides)
163777-45-5 CAPLUS
6-Quinoxalinecarboxylic acid, 3-amino-2-cyano-, ethyl ester, 1,4-dioxide

6-Quinoxalinecarboxylic acid, 3-amino-2-cyano-, ethyl ester, 1,4-dioxide (SCI) (CA INDEX NAME)

L13 ANSWER 85 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1995:189746 CAPLUS
DOCUMENT NUMBER: 122:147268
TITLE: Preparation of indenoquinoxaline derivatives for

INVENTOR (5):

Preparation of Indenoquinoxaline derivatives : electrophotographic photoreceptors Gondaira, Hideaki; Hamamoto, Isami; Nagasaki, Fumihiko; Takahashi, Hiroshi Nippon Soda Co, Japan Jpn. Kokai Tokkyo Koho, 9 pp. CODEN: JKXXAF Patent PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. APPLICATION NO. DATE DATE KIND JP 06298744 PRIORITY APPLN. INFO.: A2 19941025 JP 1993-113861 JP 1993-113861 19930416

161290-91-1 CAPLUS
11H-Indeno(1,2-b)quinoxaline-8-carboxylic acid, 11-oxo-, butyl ester (9CI)
(CA INDEX RAME)

161290-92-2 CAPLUS 11H-Indeno[1,2-b]quinoxaline-7-carboxylic acid, 11-oxo-, butyl ester (9CI) (CA INDEX NAME)

L13 ANSWER 86 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1995:267133 CAPLUS
DOCUMENT NUMBER: 123:255662
TITLE: Process for dyeing substrates with dyes containing nucleophilic and electrophilic groups and dyes for nucleophilic and electrophilic groups and dyes for SOURCE: SOURCE: Lett., UK
Brit. UK Pat. Appl., 84 pp.
CODENS AXXDU
DOCUMENT TYPE: PARKEN DE SAXXDU
PATENT INDORPATION: English
FAMILIA ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | | NO. | | | | | | | | | | | | | | | |
|-----|------|------|-----|-----|-----|-----|------|------|-----|------|------|------|-----|-----|-----|------|-----|
| | | | | | | - | | | | | | | | | - | | |
| | | 914 | | | | | | | | 3B 1 | 993- | 2356 | 0 | | 1 | 9931 | 115 |
| GB | 2272 | 914 | | | B2 | | 1996 | 0117 | | | | | | | | | |
| WO | 9412 | 717 | | | A1 | | 1994 | 0609 | 1 | NO 1 | 993- | GB23 | 44 | | 1 | 9931 | 115 |
| | W: | AT. | AU. | BB. | BG. | BR. | BY. | CA. | CH. | CZ. | DE. | DB. | DK. | DK. | ES, | FI. | GB, |
| | | HU. | JP, | KP, | KR, | KZ, | LK, | LU, | LV, | MG, | MN, | MW, | NL, | NO, | NZ, | PL, | PT, |
| | | RO. | RU. | SD. | SE. | SK. | UA. | US. | UZ. | VN | | | | | | | |
| | RW: | AT. | | | | | | | | | IE. | IT. | LU. | MC, | NL. | PT, | SE, |
| | | BF. | BJ, | CF, | CG, | CI, | CM, | GA, | GN, | ML, | MR, | NE, | SN, | TD, | TO | | |
| AU | 9454 | 311 | | | A1 | | 1994 | 0622 | - 1 | AU 1 | 994- | 5431 | 1 | | 1 | 9931 | 115 |
| | | 37 | | | | | | | | EP 1 | 993- | 9247 | 67 | | 1 | 9931 | 115 |
| EP | 6392 | 37 | | | B1 | | 1997 | 0604 | | | | | | | | | |
| | R: | AT, | | | | | ES, | FR, | GΒ, | GR, | IE, | IT, | LI, | LU, | NL, | PT. | SE |
| JP | 0850 | 3981 | | | T2 | | 1996 | 0430 | | JP 1 | 993- | 5128 | 75 | | 1 | 9931 | 115 |
| AT | 1540 | 79 | | | R | | 1997 | 0615 | 1 | AT 1 | 993- | 9247 | 67 | | 1 | 9931 | 115 |
| RS | 2102 | 690 | | | T3 | | 1997 | 0801 | 1 | ES 1 | 993- | 9247 | 67 | | 1 | 9931 | 115 |
| ZA | 9308 | 553 | | | A | | 1994 | 0720 | | ZA 1 | 993- | 8553 | | | 1 | 9931 | 116 |
| CN | 1090 | 304 | | | A | | 1994 | 0803 | | CN 1 | 993- | 1149 | 56 | | 1 | 9931 | 120 |
| US | 5474 | 580 | | | A | | 1995 | 1212 | | US 1 | 993- | 1582 | 20 | | 1 | 9931 | 129 |
| *** | 5703 | | | | | | 1997 | | | | | | | | | 9950 | 620 |

OTHER SOURCE(S): MARPAT 123:259662 US 1993-158220 A3 19931129

OTHER SOURCE(S): MARPAT 123:259662 As applying to the substrate a mixture comprising an aqueous solvent and water-soluble day which contains a nucleophilic group and an electrophilic group and heating or basifying or heating and basifying the days therefore causing mole. of the dye to join together. In this process the mol. weight of the dye increases, its water-solubility can decrease, and the affinity for textiles may be increased leading to high levels of exhaustion with good fixation and washfastness. Also claimed are polymers and oligomers of the dyes.

PAGE 1-A N- (CH2) 3-NH-

PAGE 1-B

L13 ANSMER 87 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1994:605125 CAPLUS
DOCUMENT NUMBER: 121:205125
TITLE: Preparation of [{[carboxyheterocyclyl]carbamoyl]pyrrol
idinylchiol(carbapenese as antibiotics
Jung, Frederic Henri; Arnould, Jean Claude
PATENT ASSIGNES(S): School Ltd., UK; Zeneca Pharma S.A.
SURCE: Expression CODEN: EPXXDM
DOCUMENT TYPE: PARENT: PARENT. 27 pp.

LANGUAGE: English

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INPORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|-----------|-------------------------|----------------|
| | | | | |
| EP 581500 | A1 | 19940202 | EP 1993-305607 | 19930716 |
| EP 581500 | B1 | 19980909 | | |
| R: AT, BE, CH, | DE, DK | , ES, PR, | GB, GR, IR, IT, LI, LU, | MC, NL, PT, SE |
| CA 2099818 | AA. | 19940122 | CA 1993-2099818 | 19930705 |
| AT 170859 | E | 19980915 | AT 1993-305607 | 19930716 |
| ES 2121585 | T3 | 19981201 | ES 1993-305607 | 19930716 |
| JP 06179674 | A2 | 19940628 | JP 1993-177903 | 19930719 |
| US 5441949 | A | 19950815 | US 1994-307048 | 19940916 |
| PRIORITY APPLN. INFO.: | | | EP 1992-402105 | A 19920721 |
| | | | US 1993-86836 I | 81 19930707 |
| OFFER COUNCE (C). | MADDAT | 121.20512 | 16 | |

Title compds. [I; Rl = MaCH(OH), MaCHF, CH2OH; R2,R3 = H, alkyl; Z = (iso)quinolinediyl, quinaxolinediyl, quinoxalinediyl, etc.} were prepared Thus, disodium (1R, SS, 68, 88, 2°S, 4°S)-2-(2-(8-carboxyquinol-6-ylearbampon))pyrrolidin-4-ylthiol-6-(1-hydroxyethyl)-1-methylcarbampenem-3-carboxylate, prepared in 5 steps from 6-amino-6-carboxyquinoline (preparation given), had MIC of 0.13 and 0.30 µg/ml. against Staphylococcus aureus Oxford and Escherichia coli DCO, resp. 157913-55-49 157913-56-59 157913-57-69 Preparation Research (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (Synthetic preparation of antibiotic) 157915-55-4 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dimethyl-8-nitro-, 2-propenyl ester (SCI) (CA INDEX NAME)

157915-56-5 CAPLUS 6-Quinoxalinecarboxylic acid, 8-amino-2,3-dimethyl-, 2-propenyl ester (9CI) (CA INDEX NAME)

$$H_2C = CH - CH_2 - O - C$$

$$N$$

$$Me$$

$$NHa$$

$$Me$$

157915-57-6 CAPLUS 6-Quinoxalinecarboxylic acid, 8-{[{4-(acetylthio)-1-{(2-propenyloxylcarbonyl]-2-pyrrolidinyl]carbonyl]mino]-2,3-dimethyl-, 2-propenyl ester, (28-cis)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

157915-58-7 CAPLUS
1-Azabicyclo[3.2.0]hept-2-ene-2-carboxylic acid, 6-(1-hydroxyethyl)-4-methyl-7-0xo-3-[[1-[(2-propenyloxylcarbonyl]-5-[([2,3-dimethyl-7-[(2-propenyloxylcarbonyl]-5-quinoxalinyl]amino]carbonyl]-3-pyrrolidinyl]thio]-(2-propenyl eater, $\{4R-\{3(2S^*,4S^*),4,\alpha,5\beta,6\beta(R^*)\}\}$ - [9CI) (CA INDEX RAME)

Absolute stereochemistry.

L13 ANSWER 88 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1993;490784 CAPLUS
DOCUMENT NUMBER: 119:90784
TITLE: Polycyclic compounds for cancer diagnosis and therapy

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

Tai, Seiji; Katayose, Mitsuo; Morishita, Yoshii Hitachi Chemical Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 15 pp. CODEN: JEXMAP Patent Japanese

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. KIND DATE PATENT NO. DATE JP 04288022 PRIORITY APPLN. INFO.: OTHER SOURCE(S): A2 19921013 JP 1991-49379 JP 1991-49379 MARPAT 119:90784

Polycyclic compds. I [M * H, Al, Si, P, Ga, Ge, Cd, Se, Mg, Sh, Zh; Rl-4 = H, XOM, OW, W (X = O, N, S, P, Si, CRSR6 (R5-6 = H, alkyl, aryl, aralkyl, etc.); Q -X-W linkage; W = OH, O, SH, S, etc.); k, l, m, n = 0-4; Y = halo, OR7, NR8 (R7-8 = H, (un) substituted alkyl, etc.); p = 0-2; Zl-8 enthylene, N) are respents for cancer diagnosis or therapy. Thus, Na chloroaluminonaphthalocyaninotrisulfonate (1 + 10-5M) was injected into peritonael cancer cell-bearing mice, and the treated cancer cells were sampled (isolated) and examined at 780 nm. The cancer cells were readily detected. Preparation of the compds. are given.

145946-97-2
RL: RCT (Reactant); RACT (Reactant or reagent)
(reaction of, for polycyclic cyano compound preparation for cancer diagnosis and therapy) AB

and therapy)
145964-97-2 CAPLUS
6-Quinoxalinecarboxylic scid, 2,3-dicyano-, pentyl ester (9CI) (CA INDEX NAME)

L13 ANSWER 69 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1993:191763 CAPLUS
DOCUMENT NUMBER: 188:191763
TITLE: Preparation of azamethine compounds as optical recording media
INVENTOR(S): Nagasaki, Funiniko; Hayashi, Yukio
Nagasaki, Funiniko; Hayashi, Yukio
Nippon Soda Co., Ltd., Japan
SOURCE: CODEN: JXXXAF

DOCUMENT TYPE: LANGUAGE:

JP 04288049 PRIORITY APPLN. INFO.: OTHER SOURCE(8):

KIND DATE APPLICATION NO DATE A2 19921013 JP 1991-40710 JP 1991-40710 CASRBACT 118:191763; MARPAT 118:191763 19910214 19910214

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

1.3-Bis (dicysnomethylene)-3-[(hetero)arylimino]-2,3dihydrocyclopenta[b]quinoxalines and -5,6-benxindenes [I; Y = N, CH; X =
Q-Q3; R1-R4 = H, (un]substituted alkyl, alkoxy, or NN2, halo, NO2, cyano,
OH, etc.; R5, Rs = H, (un]substituted alkyl, aryl, cycloalkyl; or RSN6
forms (hetero atom-containing) ring; R7 = H, (un]substituted alkoxy, H0, halo,
cyano, acylemino, (un]substituted alknoyloxy; R8 = H, halo,
(un)substituted alkyl, alkoxy; R9-R11 = H, alkyl] are prepared I showed
maximum sheorytion wavelengths (Amax = 750-500 nm) in a semiconductor
oscillation region, excellent solubility in organic solvents, high
sectivity.

reflectivity,
and excellent stability. Thue, 3.0 g cyclopenta[b]quinoxaline (II; Z =
H2), 2.1 g nitrosobenzene QNO (R5 = R6 = Et, R7 = OMe, R8 = H), and 70 mL
Ac20 were stirred at room temperature for 10 h to give II (Z = NQ, R5 = R6 =

R7 - OMe, R8 - N) (III) having \(\lambda\text{Max} = 825 \text{ nm. A solution of III}\) in CHC13 was spin-coated on a glass substrate and dried to form a recording medium of .appra.900 A thickness having \(\lambda\text{max} = 930 \text{ nm and } 23\) reflectively at \(\lambda\text{max} = 830 \text{ nm which formed a very clear pit by irradiation with Oa-Al-As semiconductor laser beam.

146677-81-89 146677-93-8-79
RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as intermediate for optical recording material)

146677-81-8 CAPLUS

18-Cyclopenta(b) quinoxaline-6-carboxylic acid, 1,3-bis(dicyanomethylene)-2,3-dihydro-, ethyl ceter (9CI) (CA INDEX NAME)

ΙŢ

IT

143413-59-4 CAPLUS
Benzo[a]phemazine-5-carboxylic acid, 9-(diethylamino)-, butyl ester (9CI)
(CA INDEX NAME)

L13 ANSWER 91 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1992:59404 CAPLUS
DOCUMENT NUMBER: 116:59404
ITILE: Preparation of 2,3-disubstituted quinoxalines as growth enhancers for animals
DE JONG, Anno; Puchs, Rainer
BAYENT ASSIGNEE(8): 8ayer A.-O., Germany
SOURCE: BAYEN AND PUCHS, PATENT ASSIGNEE (8): 8ayer A.-O., Germany
DOCUMENT TYPE: COUNT. PATENT AND PA

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

EP 456067 Al 19911113 EP 1991-106891 19910427
R: AT, BB, CH, DE, DK, EB, PR, GB, GR, IT, LI, LU, NL, SE

DE 4014937 Al 19911114 DE 1990-4014937 19900510

PRIORITY APPLIAL INFO::
OTHER SOURCE(8): MARPAT 116:59404

146677-98-7 CAPLUS 1H-Cyclopenta[b] quinoxaline-6-carboxylic acid, 1,1,3,3-tetrachloro-2,3-dihydro-, othyl ester (9CI) (CA INDEX NAME)

166677-62-5P
RL: SPN (Synthotic preparation); PREP (Preparation)
(preparation of, as optical recording material)
186677-62-5 CAPUS
18-Cyclopenta[b] quinoxaline-6-carboxylic acid, 1,3-bis(dicyanomethylene)-2[[4-(dicthylamino)-2-methoxyphenyl]imino]-2,3-dihydro-, ethyl ester (9CI)
(CA INDEX RAMS)

L13 ANSWER 90 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1992:532895 CAPLUS DOCUMENT NUMBER: 117:132895

117:132895

Tautomerism of 5-dicyanomethylene-9-diethylamino-5,7-dihydrobenzo[a]phenazine and ite photooxygenation to an ester in alcohol solution (Kubo, Yuji; Kuwana, Minoru; Tautsui, Sumica; Yoshida, Katsuhira
Fac. Sci., Kochi Univ., Kochi, 780, Japan
Journal of Chemical Research, Synopses (1992), (8),

AUTHOR(S): CORPORATE SOURCE .

282-3 CODEN: JRPSDC; ISSN: 0308-2342

DOCUMENT TYPE: LANGUAGE:

Title compds. [I; Rl, R2 - halo, OH, SH, alkoxy, aryloxy, alkylthio, arylthio; RlR2 = SC(X)S; X = O, S; Rl = H, halo, cyamo, CONRSR6, SOZNRSR6, SOZNRS, CH(CN)R7, OR7, COZRS; R4 = H, alkyl, halo, NO2, CONRSR6; R5 = H, alkyl, clubatituted) cytloalkyl, aryl; Re - R5, SR7; R5 = (muberituted) cytloalkyl, aryl; Re - R5, SR7; R5 = (muberituted) HANCHMOCF2, pyridine subset of the aryl) were prepared Thus, HANCHMOCF2, pyridine subset of the aryl) care prepared Thus, HANCHMOCF2, pyridine subset of the aryl aryl article were attirred 3 h in MeCN at 80° to give title compound II II to 10° tide were attirred 3 h in MeCN at 80° to give title compound II II to 10° tide were attirred 3 h in MeCN at 80° to give title compound II II to 10° tide were attirred 3 h in MeCN at 80° to give title compound II II to 10° tide were attirred 3 h in MeCN at 80° to give title compound II II to 10° tide were attirred 3 h in MeCN at 80° to give title compound II II to 10° tide were attirred 3 h in MeCN at 80° tide were attirred 3 h in MeCN attirred to 10° tide were attirred 3 h in MeCN attirred to 10° tide were attirred 3 h in MeCN attirred to 10° tide were attirred 3 h in MeCN attirred to 10° tide were attirred 3 h in MeCN attirred to 10° tide were attirred 3 h in MeCN attirred to 10° tide were attirred 3 h in MeCN attirred to 10° tide were attirred 3 h in MeCN attirred to 10° tide were attirred 3 h in MeCN attirred to 10° tide were attirred 3 h in MeCN attirred to 10° tide were attirred 3 h in MeCN attirred to 10° tide were attirred 3 h in MeCN attirred 10° tide were attirred 3 h in MeCN attirred 10° tide were attirred 3 h in MeCN attirred 10° tide were atti

138452-89-8 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-{2,2,2-trifluoro-1-methylethyl}-(9CI) (CA INDEX NAME)

138452-90-1 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(3,3,3-trifluoropropyl)- (9CI) (CA INDEX NAME)

138452-96-7 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(1-methylethyl)- (9CI) (CA INDEX NAME)

L13 ANSWER 92 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
115:222057
TITLE:
Manufacture of storage-stable dye solutions
Michne, Martin; Zillger, Hans Werner; Tegtmeyer,
Dietrich
PATENT ASSIGNEE(S):
Bayer A.-G., Germany
SOURCE:
CODEN: EPXXDM
DOCUMENT TYPE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: Patent German

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|-----|--------------------|--------|----------|-------------------|----------|
| | | | | | ***** |
| | EP 433810 | A2 | 19910626 | EP 1990-123612 | 19901208 |
| | EP 433810 | A3 | 19920115 | | |
| | R: CH, DE, FR, | GB, LI | | | |
| | DE 3942467 | A1 | 19910627 | DE 1989-3942467 | 19891222 |
| | US 5096458 | A | 19920317 | US 1990-627068 | 19901213 |
| | JP 04004263 | A2 | 19920108 | JP 1990-411097 | 19901217 |
| 210 | DETTY APPLN. INFO. | | | DR 1989-3942467 A | 19891222 |

JP 04004263 A2 19920108 JP 1990-411097 19901217
PRIORITY APPLM. INFO:
OTHER SOURCE(S):
MARPAT 115:282057

A8 Stable aqueous solns. of anionic (preferably reactive) dyes are obtained by pressure filtration of crude dye solns. in which the feed solution is obtained by stirring the optionally dried press cake or a suspension of the crude dye with a solution of Li or ammonium selts of organic or inorg. acids. Thus, 36:18 kg press cake of 1-hydroxy-2-(1,5-disulfo-2-naphthylazo)-6-(2,6-difluoro-3-chloro-4-pyrimidinylamino)-3-naphthalenesulfonic acid Na selt (1) was dissolved in 236:7 kg dt aqueous LiHCO3 solution at 45°. The composition was subjected to membrane filtration at 40 bars and 40-45°. The concentrate (94 kg) was treated with dicyandiamide 2, water 3.5, and boric acid 0.5 kg to give a stable dye solution containing 21.2% I at pH 7.5. dye solution 137682-84-9

| US 5317020 | A | 19940531 | US | 1990-610093 | | 19901105 |
|------------------------|--------|------------|----|--------------|----|----------|
| IL 111292 | A1 | 19960331 | IL | 1990-111292 | | 19901105 |
| RU 2084453 | C1 | 19970720 | RU | 1990-4831627 | | 19901105 |
| RU 2114828 | C1 | 19980710 | RU | 1993-45020 | | 19901105 |
| ZA 9008881 | A | 19910828 | ZA | 1990-8881 | | 19901106 |
| JP 03206086 | A2 | 19910909 | JP | 1990-300929 | | 19901106 |
| PL 165758 | B1 | 19950228 | PL | 1990-293823 | | 19901106 |
| PL 165854 | B1 | 19950228 | PL | 1990-293824 | | 19901106 |
| PL 166565 | B1 | 19950630 | PL | 1990-287644 | | 19901106 |
| PL 166582 | B1 | 19950630 | PL | 1990-303827 | | 19901106 |
| IL 96241 | A1 | 19960331 | IL | 1990-96241 | | 19901115 |
| LV 10713 | B | 19951020 | LV | 1993-142 | | 19930225 |
| US 5686609 | A | 19971111 | US | 1994-208672 | | 19940311 |
| AU 9459245 | A1 | 19940603 | AU | 1994-59245 | | 19940331 |
| AU 668018 | B2 | 19960418 | | | | |
| NO 9500239 | A | 19910507 | NO | 1995-239 | | 19950123 |
| NO 180193 | B | 19961125 | | | | |
| NO 180193 | С | 19970305 | | | | |
| NO 9500240 | A | 19910507 | NO | 1995-240 | | 19950123 |
| NO 179580 | В | 19960729 | | | | |
| NO 179580 | c | 19961106 | | | | |
| US 5618938 | A | 19970408 | UŞ | 1995-479634 | | 19950607 |
| FI 9502956 | A | 19950615 | FI | 1995-2956 | | 19950615 |
| FI 9502957 | A | 19950615 | FI | 1995-2957 | | 19950615 |
| FI 9800227 | A | 19980202 | FI | 1998-227 | | 19980202 |
| PRIORITY APPLN. INFO.: | | | FR | 1989-14517 | A | 19891106 |
| | | | FR | 1990-7534 | A | 19900615 |
| | | | FI | 1990-5444 | A | 19901102 |
| | | | NO | 1990-4802 | A | 19901105 |
| • | | | US | 1990-610093 | A3 | 19901105 |
| | | | IL | 1990-96241 | A3 | 19901115 |
| | | | US | 1994-208672 | A3 | 19940311 |
| | | | FI | 1995-2956 | A | 19950615 |
| OTHER SOURCE(S): | MARPAT | 115:279818 | | | | |
| | | | | | | |

The title compds, I [m = 1-3; Ar, Ar' = thienyl, (substituted) Ph, stc.; X = H; X' = H, ON; or XX' = oxo, dislkylaminoslkyloxyimino, etc.; Y = N, CX'; X'' = Hor X'X'' = carbon-carbon bond; O = H, slkyl, (CR12)Am'; q = 2 or 3; Am' = piperidino, 4-bensylpiperidino, etc.; R = H, Me, (CR12)hL; n = 2-6; L = H, amino; T = CO, (ON)H; N = O, S; Z = H, M, or ON When T = CO; or Z = M when T = C(W)NH; M = A, slkyl, (substituted) phenylalkyl, etc.] were prepared I are neurokinin and substance P antagonists (no data). Reaction of amine II (21 = H) with 2,4-dichlorobensoyl chloride in the presence of EllN gave II (21 = 2,4-dichlorobensoyl) isolated as its HCl salt. I are also useful as allergy and inflammation inhibitors (no data). 135956-087 135956-48-8P

RL: USES (Uses)
(dye, storage-stable aqueous solns. containing)
137623-64-9 CAPLUS
2-Anthracenssulfonic acid, 1-amino-4-[[4-[[(2,3-dichloro-6-quinoxaliny]) carbonyl]methylamino]methyl]-2-sulfophenyl]amino]-9,10-dihydro-9,10-dioxo-(9CI) (CA INDEX NAME)

L13 ANSWER 93 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1991:679818 CAPLUS DOCUMENT NUMBER: 115:279818

115:279818
Preparation of piperidine derivatives as neurokinin and substance P antagonists
Emonds-Alt, Xevier; Goulaouic, Pierre; Proietto, Vincenzo; Van Broeck, Didder SANOFI, Fr.
Eur. Pat. Appl., 84 pp.
CODEN: EPIXON
Patent
Prench TITLE:

INVENTOR (8):

PATENT ASSIGNEE (S): SOURCE:

DOCUMENT TYPE: LANGUAGE:

PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. DATE KIND A2 A3 DE, DK, A1 B1 A1 EP 428434 EP 428434 R: AT, BE, CH, FR 2654100 19910522 19911009 ES, FR, 19910510 EP 1990-403125 19901106 GB, GR, IT, LI, LU, NL, FR 1989-14517 FR 2654100 FR 2654100 FR 2663329 FR 2663329 FI 97540 FI 97540 CA 2029275 NO 9004802 NO 177299 NO 177299 NO 9065838 19891106 19920221 19920221 19911220 19921016 19960930 19970110 19910507 19910507 19950515 19950823 FR 1990-7534 19900615 PI 1990-5444 19901102 CA 1990-2029275 NO 1990-4802 19901105 19901105 AU 9065838 AU 649973 HU 56543 19910523 19940609 19910930

AU 1990-65838

HU 1990-7027

19901105

19901105

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation): THU (Therapeutic use); BIOL (Biological study): RRED (Preparation): USES (Uses) (preparation) of, as neurokinin antagonist) 15956-48-8 CAPLUS 6-Quinoxalinecarboxamide, N-[2-03,4-dichlorophenyl)-4-[4-(phenylmethyl)-1-piperidnyl]butyl]-, monohydrochloride (9CI) (CA INDEX NAME)

• HCl

L13 ANSWER 94 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION MUMBER: 1991:492302 CAPLUS
DOCUMENT NUMBER: 115:92302 CAPLUS
TITLE: Properation of benzo[a]phenazine derivatives
Shirai, Hiroyoshi; Hanabusa, Kenji; Oce, Okikezu; Uda, Yoshihiro
PATENT ASSIGNEE(S): Taisho Pharmaceutical Co., Ltd., Japan
JDN. Kokai Tokkyo Koho, 3 pp.
CODEN: JKKXAF
DOCUMENT TYPE: Patent

Patent Japanese 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|-----------|-----------------|----------|
| | | | | |
| JP 03066698 | A2 | 19910322 | JP 1989-204211 | 19890807 |
| PRIORITY APPLN. INFO.: | | | JP 1989-204211 | 19890807 |
| OTHER SOURCE(S): | MARPAT | 115:92302 | | |
| ar | | | | |

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The title derive. I (R = Q1-Q3), useful as antitumor agents (no data), are prepared Thus, 1.90 g I (R = H) was stirred with 1.75 g II in DMF in the presence of powdered K2CO3 at 50-60° for 24 h to give 1.14 g I (R = Q1).

01).
13442-59-8P 135412-60-1P 135438-74-3P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SN (Synthetic preparation); THU (Therapoutic use); IT

BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as antitumor agent) 135412-59-5 CAPLUS and 135412-59-5 CAPLUS and 135412-59-5 CAPLUS and 12-0-(1-methylethylidene)-, 6-[6-[[2-(dimethylamino) ethyl] hamino| carboxyl-10-methoxybenso [a] phenazine-9-carboxylate], monohydrochloride (9CI) (CA INDEX NAME)

Absolute stereochemistry.

• HC1

135412-60-1 CAPLUS

α-D-Glucopyranose, 5-[6-[[[2-(dimethylamino)ethyl]amino]carbonyl]-5hydroxy-10-methoxybenzo[a]phenazine-9-carboxylate], monohydrochloride
(9CI) (CA INDEX NAME)

Absolute stereochemistry.

• HC1

135438-74-3 CAPLUS

α-D-Glucofuranose, 1,2:3,5-bis-O-(1-methylethylidene)-,
6-[[[2-(dimethylamino|ethyl)amino|carbonyl]-5-hydroxy-10methoxybenzo[s]phenazine-9-carboxylate (9CI) (CA INDEX NAME)

Absolute Stereochemistry.

L13 ANSWER 95 OF 161 CAPLUS COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER: 1991:420992 CAPLUS
DOCUMENT NUMBER: 115:20992
TITLE: Substituted tetraquinoxalinoporp

Substituted tetraquinoxalinoporphyrazine derivative with near-infrared absorption Nagasaki, Fumihiko; Hatano, Hirosi; Takahashi, Hiroshi Nippon Soda Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 5 pp. CODEN: JKXXAF Patent Japanose

INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. JP 02289575
PRIORITY APPLM. INFO.:
OTHER SOURCE(S):
GI

A2 19901129 APPLICATION NO.

JP 1989-73155
JP 1989-32144 A MARPAT 115:20992

STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The derivative is I $\{Z=H;R\}$ -4 = halo, (substituted) alkyl, alkoxy, alkylthio, phenylthio, phenylthio, phenyloxy, and seter, carboxyl, amide, amino; k, l, m, n = 0.4; k + 1 + m + n \geq 1; H = 2H, metal metal oxide, setal hydroxide, acyl metal, metal alkoxide, metal siloxide, metal halide). The derivative, with high near-IR absorption and solubility to an organic solvent,

IT

useful for an optical recording medium, electrophotog., photoreceptore, radox catalysts, flower preservatives, etc.
11438-14-2-134382-43-7 134382-44-8
114382-45-9 134382-45-0
RL: RCT (Reactant): RACT (Reactant or reagent)
(near-IR-absorbing)
13438-14-5 CAPLUS
Copper, [tetracthyl 37H.39H-tetracquinoxalino[2,3-b:2',3'-g:2'',3''-1:2'',3''-q]porphyraxine-2,11.70,29-tetracarboxylato(2-)N37,N38,N39,N40]-, (SP-4-1)- (9CI) (CA INDEX NAME)

PAGE 3-A

134382-43-7 CAPLMS
Tin, dichloro[tetraethyl 37H,39H-tetraquinoxalino[2,3-b:2',3'-g:2'',3''-1;2'',3''-q]porphyraxine-2,11,20,29-tetracarboxylato(2-)-N37,N38,N39,N40]-, (OC-6-12)- (9CI) (CA INDEX NAME)

PAGE 1-A

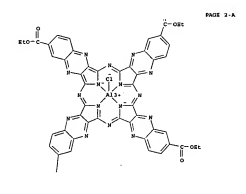
PAGE 3-A

RN 13438-44-8 CAPLUS
CN Vanadium, cxo(tetraethyl 37H,39B-tetraquinoxalino[2,3-b:2',3'-g:2'',3''-l:2'',3''-g)porphyrazine-2,11,20,29-tetracarboxylato(2-)-H37,N38,N39,N40]-, (8P-5-12)- (9C1) (CA INDEX NAMB)

PAGE 3-A

134382-45-9 CAPLUS
Aluminum, chloro[tetraethyl 37H,39H-tetraquinoxalino[2,3-b:2',3'-g:2'',3''1:2'',3''-d]porphyrazine-2,11,20,29-tetracarboxylato(2-)N37,N38,N39,N40]-, (SP-5-12)- (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 3-A

RN 134382-46-0 CAPLUS
CN Zinc, [tetraethyl 37H,39H-tetraquinoxalino[2,3-b;2',3'-g;2'',3''-li2'',3''-d] porphyrazine-2,11,20,29-tetracarboxylato(2-)-N37,N38,N39,N40]-, (SP-4-1)- (9CI) (CA INDEX NAME)

LI3 ANSWER 96 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1991:104337 CAPLUS
DOCUMENT NUMBER: 114:104337
TITLE: Rectrochemical investigations on the analysis of reactive dyes with monoszo- and monoanthraquinone structures

Eto-C

STRUCTURES

HOR(8): Sahm, Uwe; Knittel, Dierk; Schollmeyer, Eckhard

DORATE SOURCE: Dech. Textilforechungszent. Nord-West e.V., Krefeld,

W-4150/1, Germany

RCE: Freeenius' Journal of Analytical Chemistry (1990),

338(7), 834-30

CODEN: FJACES; ISSN: 0937-0633

JOURNAT TYPE: Journal

JOURNAT TYPE: Journal

JOURNAT TYPE: Journal

Qual. voltammetric determination of reactive dyes is possible down to concns. AUTHOR(S): CORPORATE SOURCE:

DOCUMENT TYPE: LANGUAGE:

.apprx.10-5 mol/L using d.c. polarog. Using rotating solid electrodes (glassy carbon) a detection limit of .apprx.10-4 mol/L can be obtained. Reduction involves chiefly only the azo- and anthraquincid groups. Discrimination between an original reactive dye and its hydrolysate (containing a hydrolytically destroyed reactive group) can be achieved, if the dye contains further cleavable groups such as benxamido substituents in conjugation to the chromophore as is seen with an azo red dye. Cleavage of such a group shifts the reduction potential for .apprx.130 mV to more neg. values.

values.
132164-73-5
RL: ANT (Analyte); ANST (Analytical study)
(determination of, voltammetric, electrochem. reactions in)
132366-73-5 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-{[4-[{(2,3-dichloro-6-quinoxalinyl)-arthylmino]methyl]-3-sulfophenyl]amino]-9,10-dihydro-9,10-dioxo- (9CI) (CA INDEX NAME)

PAGE 2-A

L13 ANSWER 97 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
1990:611948 CAPLUS
TITLE:
POOLEMENT NUMBER:
113:211948
POLENTIAL ALTHOR(S):
AUTHOR(S):
CORPORATE SOURCE:
SOURCE:
CORPORATE SOURCE:
DOCUMENT TYPE:
LINGUAGE:
DOCUMENT TYPE:
LINGUAGE:
OTHER SOURCE(S):
GT APPLIES OF THE SOURCE SIGNED ACCES ASSESSED ACCES ACCESSED ACCESSE

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI

Ring analogs and derivs. of antimitotic antitumor 1,2-dihydropyrido[3,4-b]pyrazinylcarbamates, e.g., I, were prepared form benzoic acids, e.g. 4,3,5-C(020N)2C6H20C3N, and pyridylcarbamates, e.g. II. In vitro evaluation indicated that activity was reduced by removal of the pyridine ring nitrogen of I and destroyed by increasing the basicity of the pyrazine ring of I as in the case of aminopyridotriazinylcarbamates III. 130145-39-0P
RL: SPN (Synthetic preparation); PREP (Preparation) (preparation, borohydride reduction and neoplasm inhibiting activity of) 130145-39-0 CAPLUS (-Quinozilinecarboxylic acid, 8-amino-2-phenyl-, ethyl ester (9CI) (CA INDEX NAME)

L13 ANSWER 98 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1990:179846 CAPLUS
DOCUMENT NUMBER: 112:179846
TITLE: Synthesis and antimicrobial activity of some new

AUTHOR (s):

CORPORATE SOURCE:

2,3-dichloroquinoxaline-6-sulfonyl amino acid and dipeptide derivatives Kora, P. A.; Hussein, M. E.; El-Sayed, R. A.; El-Naggar, Ahmed M. Fac. Sci., Al-Athar Univ., Naer, Egypt Polieh Journel of Chemietry (1988), 62(7-12), 749-56 CDDEN: PJCHDQ; ISSN: 0137-5083 Journal Snglish CAGREACT 112:179846 SOURCE:

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S):

Title amino acid derivs. I (X = Gly, Ala, Val, Leu, DL-Leu, Ser, DL-Ser, Phe, Tyr; R = OH) were prepared by sulfonylating the corresponding amino acids with sulfonyl chloride II in the presence of St3N. Me esters I (X = Gly, DL-Ala, Leu, Ser; R = OMe) were prepared similarly from II and the appropriate amino acid Me ester hydrochlorides. The above Me esters were converted into hydrazides I (X = Gly, DL-Ala, Leu, Ser, R = NINN3). Dispetides III (X1-X2 = Phe, DL-Ala, Phe-Leu, Tyr-Gly, Tyr-DL-Ala, Tyr-Leu) were prepared by the DCC method. All the above compds. were active against a number of microorganisms.

1/1715-80-78 171735-80-89 117135-90-19
117135-80-79 117135-90-91 117135-90-19
117135-90-99 117136-00-69 117136-01-79
117136-03-19
RL BAC (Sicological activity or effector, except adverse); BSU (Biological study), unclassified), SPN (Synthetic preparation); BIOL (Biological study); PRPO (Preparation)
(preparation and antimicrobial activity of)
Glycine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl)- (SCI) (CA INDEX NAME)

117195-86-5 CAPLUS L-Alanine, N-{{2,3-dichloro-6-quinoxalinyl}eulfonyl}- (9CI) (CA INDEX NAME)

Absolute stereochemistry

117195-87-6 CAPLUS L-Valine, N-{(2,3-dichloro-6-quinoxalinyl)sulfonyl}- (9CI) (CA INDEX RAME)

Absolute stereochemistry.

117195-88-7 CAPLUS L-Leucine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 117195-89-8 CAPLUS
CN Leucine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]- (9CI) (CA INDEX NAME)

RN 117195-90-1 CAPLUS CN L-Serine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

117195-91-2 CAPLUS Serine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]- (9CI) (CA INDEX NAME)

117195-97-8 CAPLUS Glycine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-, hydrazide (9CI) (CA INDEX NAME)

117195-98-9 CAPLUS
Alenine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-, hydraxide (9CI) (CA
INDEX NAME)

117195-99-0 CAPLUS L-Leucine, N-[4:3-dichloro-6-quinoxalinyl)eulfonyl]-, hydrazide (9CI) (CA INDEX RAME)

Absolute stereochemistry

117196-00-6 CAPLUS L-Serine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-, hydrazide (9CI) (CA INDEX NAME)

Absolute stereochemistry.

117196-01-7 CAPLUS
Alanine, N-[N-[(2,3-dichloro-6-quinoxalinyl) sulfonyl]-L-phenylalanyl]-,
methyl ester (9CI) (CA INDEX RAME)

117196-02-8 CAPLUS L-Leucine, N. [N-[(2.3-dichloro-6-quinoxalinyl)sulfonyl]-L-phenylalanyl]-, methyl ester (SCI) (CA INDEX NAME)

117196-03-9 CAPLUS Glycine, N-[N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-L-tyrosyl]-, methyl ester (9C1) (CA INDEX NAME)

117196-04-0 CAPLUS
Alenine, N-[N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-L-tyrosyl]-, methyl ester (9C1) (CA INDEX NAME)

117196-05-1 CAPLUS L-Leucine, N-[N-[(2,3-dichloro-6-quinoxaliny1)sulfony1]-L-tyrosy1)-, methyl ester (9CI) (CA INDEX NAME)

117195-93-4P 117195-94-5P 117195-95-6P 117195-96-7P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation, hydrazinolysis, and antimicrobial activity of) 117195-93-4 CAPUUS Olycine, No. 1(2.3-dichloro-6-quinoxalinyl)sulfonyl)-, methyl ester (9CI) (CA INDEX NAME)

Alanine, N=[(2,3-dichloro-6-quinoxalinyl)sulfonyl}-, methyl ester (9CI) (CA NDSX NAME) 117195-94-5 CAPLUS

117195-95-6 CAPLUS L-Leucine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-, methyl ester (9CI) (CA INDEX NUME)

Absolute stereochemistry.

117195-96-7 CAPLUS L-Serine, N-[(2,3-dichloro-6-quinoxelinyl)sulfonyl]-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

117195-92-3P 117222-08-9P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation, peptide coupling reaction, and antimicrobial activity of)
117195-92-3 CAPLUS
L-Tyrosine, N-[(2,3-dichloro-6-quinoxaliny1)sulfony1]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

concentration in blood as conventional emulsions with large particle size. NC-239 (I, R = Bu) (28 mg) was dispersed in 4 g Panacete 810 (triglyceride), homogenized with 800 mg Nikkol TO-10M [poly(oxyethylene) sorbitan fatty acid ester), 880 mg glycerin, and H30 to 40 mL (pH 7.4), charged into ampuls, and sterilized to give an emulsion (average particle size 40 mm), which was i.v. administered to mice bearing lung cancer at 25 mg (as NC-239)/kg/day for 8 days to show T/C (treated group/control group) survival rate >2184, vs. 1554, for an emulsion with 250 nm average particle

eurvival rate xair, --size.
106224-68-4 106225-12-1 106225-21-2
RI. BIOL (Biological study)
(antitumor emulsions containing, with improved bioavailability)
166224-68-4 CAPLUS
Benzo[a]phenazine-9-carboxylic acid, 6-{[{2-(dimethylamino)ethyl]amino]carbonyl]-5-hydroxy-10-methoxy-, butyl ester (9CI) (CA INDEX NAME)

106225-12-1 CAPLUS Benzo (alphenazine-9-carboxylic acid, 6-{[[2-(dimethylamino)ethyl]amino]car bonyll-5-hydroxy-10-methoxy-, propyl ester (SCI) (CA INDEX NAMS)

106225-21-2 CAPLUS

Benzo(a|phenazine-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]carboxyl-1-5-hydroxy-10-methoxy-, decyl ester (9CI) (CA INDEX NAME)

L13 ANSWER 100 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN

117222-08-9 CAPLUS L-Phenylalanine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]- (9CI) (CA INDEX NAMB)

Absolute stereochemistry.

L13 ANSWER 99 OP 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
117:25680
Bloavailability-improved anticancer emulsions
containing benzo[a]phenazines
Yameguchi, Hiroshi, Ozawa, Yasuo; Kano, Akira;
Heyashi, Hidefumi; Shoji, Minoru; Athara, Hirokazu;
Kotomo, Susumu; Nekaike, Shiro
PATEMT ASSIGNEE(S):
50URCE:
Jon. Kokai Tokkyo Koho, 3 pp.
CODEN: JON. XXXAF
DOCUMENT TYPS:
PAMILUT ACC. NUM. COUNT:
1
Japanese
PAMILUT ACC. NUM. COUNT:
1

A2 19890606 B4 19940817

PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

KIND DATE PATENT NO.

APPLICATION NO. DATE JP 1987-304047 JP 1987-304047 19871201

JP 01143834 JP 06062418 PRIORITY APPLN. INFO.: OTHER SOURCE(S): GI MARPAT 112:25680

Anticancer emulsions contain benzo(a)phenazines (I; R = alkyl) with average particle size 40-70 nm. The emulsions do not show sharp decrease of I

ACCESSION NUMBER:

1989:116960 CAPLUS
110:116980
Reactive dichloroquinoxaline group-containing dyes
Jasger, Horat; Stochr, Frank Michael; Herd, Karl
Josef; Henk, Hermann; Schwarz, Max; Koecher, Juergen
Bayer A.-O., Fed. Rep. Ger.
CODEN; GHXXBX
PAtent
German
1
1 TITLE: INVENTOR(S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------|--------|----------|-----------------|----------|
| PATENT NO. | ****** | | ALI DI GILI IO. | |
| | | | | |
| DE 3707549 | A1 | 19880922 | DE 1987-3707549 | 19870310 |
| EP 281898 | A2 | 19880914 | EP 1988-103052 | 19880301 |
| EP 261898 | A3 | 19890111 | | |
| EP 261898 | B1 | 19910710 | | |

R: CH, DE, FR, GB, LI
PRIORITY APPLIN. INFO.:
OTHER SOURCE(S): DE 1987-3707549 A 19870310 CASREACT 110:136960; MARPAT 110:136960

The title dyes (XOISD1)1-3GDN(R)Z, [D. D1 = direct bond, aromatic carbocyclic bridging groups. aromatic haterocyclic bridging group. G = chromophore residue; R = N, (un) substituted (1-4 alkyl; X = CH.CHZ, CHZCHZY; Y = alkali-cleavable substituted; Z = fiber-reactive residue], useful for dyeing or printing RO or carbonanide group-containing materials, are prepared i-Amino-8-hydroxy-3, 6-naphthalenedisulfonic acid was condensed with 2,3-dichloroquinoxaline-6-carbonyl chloride, and the condensate coupled with diazotized 2-amino-6-sulfatoethylsulfonyl-1-naphthalenesulfonic acid, producing I, which dyad cotton in a fast bluish-red shade.

119385-60-3P
RL: PREP (Preparation)
(manufacture of, as reactive blue dye)
119385-60-31 CADLUS
6-Quinoxalinecarboxamide, N,N'-[[6,13-dichloro-4,11-bis[[2-(sulfoxylethyl]sulfonyl]-3.10-triphenodioxazinediyl]bis(mino-2,1-ethanediyl)]bis[2,3-dichloro-N-[2-(sulfoxy)ethyl]sulfonyl) -3.10-triphenodioxazinediyl]bis(mino-2,1-ethanediyl)]bis[2,3-dichloro-N-[2-(sulfoxy)ethyl]- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{MO}_3\text{SO}-\text{CH}_2-\text{CH}_2 & \text{M} \\ \text{C1} & \text{N} & \text{O} & \text{HO}_3\text{SO}-\text{CH}_2-\text{CH}_2-\text{S} = 0 \\ \text{C1} & \text{N} & \text{C1} & \text{O} & \text{C1} \\ \text{C1} & \text{N} & \text{C1} & \text{O} & \text{C1} \\ \end{array}$$

119385-47-6P
RL: PREP (Preparation)
(manufacture of, as reactive yellow dye)
119385-47-6 CAPUS
3-Pyridinemechanesulfonic acid, 1-[2-[((2,3-dichloro-6-quinoxalinyl)earbonyl]amino]ethyl]-1,2-dihydro-6-hydroxy-4-methyl-2-oxo-5-[(1-sulfo-6-[(2-(sulfooxy)ethyl]sulfonyl)-2-naphthalenyl]azo]- (9CI) (CA
INDEX NUME)

L13 ANSWER 101 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1989:23865 CAPLUS DOCUMENT NUMBER: 110:23865

106224-81-1P 106224-87-7P 106224-90-2P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as intermediate for antitumor agents) 106224-71-9 CAPLIS Banzo[a]phenazine-6,9-dicarboxylic acid, 10-ethoxy-5-hydroxy-, diethyl ester (9CI) (CA INDEX NAME)

106224-72-0 CAPLUS Benzo[a]phenazin-6,9-dicarboxylic acid, 10-ethoxy-5-hydroxy-, 9-butyl 6-ethyl ester (9C1) (CA INDEX NAME)

106224-74-2 CAPLUS Benzo[a]phenazin-6,9-dicarboxylic acid, 5-hydroxy-10-propoxy-, diethyl ester (9CI) (CA INDEX NAME)

106224-75-3 CAPLUS
Benzo[a]phenezine-6,9-dicarboxylic acid, 5-hydroxy-10-propoxy-, 6-ethyl
9-propyl ester (9CI) (CA INDEX NAME)

INVENTOR(S):

Preparation of 5-hydroxybenzo[a]phenazine-6-carboxylates as intermediates for antituor sgents Uda, Yoshihiro; Kumazawa, Yukinari; Nakagami, Yoji; Amano, Takshiro; Soda, Kaoru; Sakakibara, Nisaku Taisho Pharmaceutical Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 8 pp. CODEN: JOKUAF

PATENT ASSIGNEE(S):

Patent Japanese 1 DOCUMENT TYPE:

APPLICATION NO. PATENT NO. KIND DATE DATE JP 63083073 JP 06076393 PRIORITY APPLN. INFO.: OTHER SOURCE(S): GI 19880413 19940928 JP 1986-229104 19860927 JP 1986-229104

Title compds. I (R1 = H, halo, Me, OH, alkoxy; R2 = CO2R4, CONRSR6; R3 = alkyl; R4 = H, alkyl, cycloalkyl, PhCH2, Ph; R5, R6 = H, alkyl; RSR6N = heterocyclyl) are prepared as intermediates for benzo(a)phenazina-6-carboxamide antitumor agents. Treatment of Et 3-hydroxy-1,4-dihydro-1,4-dioxo-2-naphthoate with CHCO2Et in THF in the presence of RtN gave Et 3-ethoxycarbomyloxy-1,4-dihydro-1,4-dioxo-2-naphthoate, followed by cyclocondensation with Me 4,5-diamino-2-methoxybenzoate in DNF gave I (R1 MeO, R2 = MeO2R, R3 = RC1), which was refluxed with Me2N(CH2)2NH2 in C6H6 to afford the corresponding amide.

IT

106224-71-99 106224-72-09 106224-74-29 106224-75-39 106224-76-49 106224-78-69

106224-76-4 CAPLUS Benzo[a]phenezine5, 9-dicarboxylic acid, 5-hydroxy-10-propoxy-, 9-butyl 6-ethyl ester (9C1) (CA INDEX NAME)

106224-78-6 CAPLUS Benzo[a]phenazine-6,9-dicarboxylic acid, 10-butoxy-5-hydroxy-, 9-butyl 6-ethyl ester (9CI) (CA INDEX NAME)

106224-81-1 CAPLUS Benzo[a]phenaxine-6,9-dicarboxylic acid, 5-hydroxy-, diethyl ester (9CI) (CA INDEX NAME)

106224-87-7 CAPLUS

Benzo[a]phenazine-6-cerboxylic acid, 9-[(ethylamino)carbonyl]-5-hydroxy-10-methoxy-, ethyl ester (9CI) (CA INDEX NAME)

106224-90-2 CAPLUS Benzo[a]phenazine-6,9-dicarboxylic acid, 5-hydroxy-10-metho 9-(2,2-dimethylpropyl) 6-ethyl ester (9CI) (CA INDEX NAME)

L13 ANSWER 102 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1988:590818 CAPLUS DOCUMENT NUMBER: 109:190818

109:190818
Synthesis and antimicrobial activity of some new
2,3-dichloroquinoxaline-6-sulfonyl amino acid and
dipeptide derivatives
Kora, P. A.; Hussein, M. E.; El-Sayed, R. A.;
El-Naggar, A. M.
Fac. Sci., Al-Azhar Univ., Nasr, Egypt
Journal of the Serbian Chemical Society (1987), 52(9),
529-35
CODEN: JSCSEN; ISSN: 0352-5139
Journal

AUTHOR(S):

CORPORATE SOURCE:

Journal English CASREACT 109:190818

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): G1

Title amino acide I (X = Gly, Ala, Val, Leu, DL-Leu, Ser, DL-Ser, Phe, Tyr; R = OH) and Me ceters I (X = Gly, DL-Ala, Leu, Ser; R = OMe) were prepared by treating sulfonyl chloride II with the appropriate amino acids and amino acid Me esters. Hydratides I (X = Gly, DL-Ala, Leu, Ser; R = NHNM2) were prepared by treating the corresponding Me esters with NHZNH2. Dispetide Me esters I (X = Phe-DL-Ala, Phe-Leu, Tyr-Gly, Tyr-DL-Ala,

Tyr-Leu; R = ONe) were also prepared The above compds, were active against a number of microorganisms.

117:195-187-49 117:195-86-59 117:195-87-59
117:195-88-79 117:195-98-89 117:195-90-19
117:195-97-29 117:195-90-19
117:195-97-09 117:196-00-69 117:196-01-79
117:196-03-199 117:196-01-99 117:196-01-09
117:196-03-199 117:196-01-09 117:196-01-09
117:196-05-19
RL: BAC (Biological activity or effector, except adverse); BSU (Biological actudy, unclassified); SPN (Synthetic preparation); BIOL (Biological actudy); PREP (Preparation)
(preparation and antimicrobial activity of)
117:195-95-4 CAPLUS
Clycine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]- (9CI) (CA INDEX NAME)

117195-86-5 CAPLUS L-Alanine, N-[{2,3-dichloro-6-quinoxalinyl)sulfonyl}- (9CI) (CA INDEX NAME)

117195-87-6 CAPLUS L-Valine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

117195-88-7 CAPLUS L-Leucine, N-{(2,3-dichloro-6-quinoxalinyl)eulfonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

117195-89-8 CAPLUS Leucine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]- (9CI) (CA INDEX NAME)

117195-90-1 CAPLUS L-Serine, N-[{2,3-dichloro-6-quinoxalinyl}sulfonyl]- (9C1) (CA INDEX NAME)

Absolute stereochemistry

117195-91-2 CAPLUS
Serine, N-{(2,3-dichloro-6-quinoxalinyl)sulfonyl}- (9CI) (CA INDEX NAME)

117195-97-8 CAPLUS Clycine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-, hydrazide (9CI) (CA INDEX NAME)

117195-98-9 CAPLUS Alanine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-, hydrazide (9CI) (CA INDEX NAME)

117195-99-0 CAPLUS L-Leucine, N- ((2,3-dichloro-6-quinoxalinyl)sulfonyl]-, hydrezide (9CI) (CA INDEX NAME)

117196-00-6 CAPLUS L-Serine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-, hydrazide (9CI) (CA INDEX NAME)

117196-01-7 CAPLUS
Alanine, N-[N-[(2,3-dichloro-6-quinoxaliny1)sulfony1]-L-phenylelany1]-,
methyl seter (9CI) (CA INDEX NAMS)

RN 117196-02-8 CAPLUS
CN L-Leucine, N-[N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-L-phenylalanyl].

methyl ester (9CI) (CA INDEX NAME)

117196-03-9 CAPLUS
Glycine, N-[N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl}-L-tyrosyl]-, methyl
ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

117196-04-0 CAPLUS
Alanine, N-[N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-L-tyrosyl]-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

117196-05-1 CAPLUS L-Leucine. N. [N-{(2,3-dichloro-6-quinoxalinyl)sulfonyl]-L-tyrosyl]-, methyl ester (9C1) (CA INDEX NAME)

117195-93-4P 117195-94-5P 117195-95-6P 117195-96-7P 117195-96-7P RE: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation, hydrazinolysis, and antimicrobial activity of) 117195-93-4 CAPUUS Glycine, N. (2,3-dichloro-6-quinoxalinyl)sulfonyl)-, methyl ester (9CI) (CA INDEX NAME)

117195-94-5 CAPLUS
Alanine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl]-, methyl ester (9CI)
(CA INDEX NAME)

117195-95-6 CAPLUS L-Leucine, N-[(1,3-)-dichloro-6-quinoxalinyl)sulfonyl]-, methyl ester (9CI) (CA INDEX NAME)

117195-96-7 CAPLUS L-Serine, N-[(2,3-dichloro-6-quinoxalinyl)sulfonyl}-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

117195-92-3F 117222-08-99
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation, peptide coupling, and antimicrobial activity of)
17135-92-3 CAPLUS
L-Tyrosine, N-[(2,3-dichloro-6-quinoxaliny1)sulfony1]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

117222-08-9 CAPLUS L-Phenylelanine, N-{(2,3-dichloro-6-quinoxalinyl)sulfonyl]- (9CI) (CA INDEX NAME)

DOCUMENT TYPE: LANGUAGE:

L13 ANSMER 103 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1988:18518 CAPLUS
TITLE: Synthesis and biological activity of some new
2.3-dihydroxyquinoxaline-6-sulfonyl maino acids and
dipeptide derivatives P. A.; El-Sayed, R. A.
CORPORATE SOURCE: 81-Naggar, A. H.; Kora, P. A.; El-Sayed, R. A.
Fac. Sci., Al-Azhar Univ., Ceiro, Egypt
Journal of the Serbian Chemical Society (1986),
51(9-10), 441-7
CODEN: JSCSEN; ISSN: 0352-5139
JOURNET TYPE:

Title amino acid derivs. I (X = Ale, DL-Ala, Val, DL-Val, Leu, Phe, etc.) were prepared by sulfonylating the corresponding amino acid with sulfonyl chloride II. Some of the above amino acid derivs. were converted into their Ne caters and hydrazides. I (X = Val, Leu, Phe, Tyr) were coupled with amino acid Me esters by the DCC method to give the corresponding dipeptide derivs. All of the above synthesized derivs, were active against a number of microorganisms, e.g., Bacillus cerues and Candida utilis. 11259-97-89 112159-99-99 112170-00-99 112170-01-1P 112170-02-2P 112170-03-1P 112170-04-99 112170-04-91 12170-04-91 12170-04-91 12170-04-91 12170-06-89 112170-19-1P 112170-26-09
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SNN (Synthetic preparation); BIOL (Biological study); PREP (Preparation) (preparation and antimicrobial activity of) 112169-97-8 CAPUUS L-beucine, N-IM-(1(2,2),4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]-L-tyrosyl]-, methyl ester (9CI) (CA INDEX NAME)

112169-98-9 CAPLUS L-Leucine, N. [A. [1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]-L-phenylalanyl]-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

112169-99-0 CAPLUS L-Leucine, N-[N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]-L-leucyl]-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

112170-00-0 CAPLUS L-Leucine, N.[H. ([1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]-L-valyl)-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

112170-01-1 CAPLUS Alanine, N-(N-(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl}-L-valyl}-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

112170-05-5 CAPLUS
Valime, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]-, hydrazide (9C1) (CA INDEX NAME)

112170-06-6 CAPLUS L-Valine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]-, hydraxide (9CI) (CA INDEX NAME)

Absolute stereochemistry.

112170-07-7 CAPLUS
Alanine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxelinyl)sulfonyl]-,
hydrazide (9CI) (CA INDEX NAME)

112170-08-8 CAPLUS L-Alanine, N-{(1, 2, 3, 4-tetrahydro-2, 3-dioxo-6-quinoxaliny1) sulfony1}-, hydrazide (SCI) (CA INDEX NAME)

Absolute stereochemistry.

112170-02-2 CAPLUS L-Phenylalanine, N-{(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl}-, hydraide (SCI) (CA INDEX NAME)

Absolute stereochemistry.

112170-03-3 CAPLUS Leucine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxaliny1)sulfony1]-, hydrazide (9C1) (CA INDEX NAMS)

112170-04-4 CAPLUS L-Leucine, N-{(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl}-, hydraxide (9C1) (CA INDEX NAME)

Absolute stereochemistry.

112170-18-0 CAPLUS Serine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]- (9CI) (CA INDEX NAME)

112170-19-1 CAPLUS L-Serine, B- (1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl)- (9CI) (CA INDEX NAME)

112170-26-0 CAPLUS
Glycine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]- (9CI)
(CA INDEX NAME)

112170-20-4P 112170-22-6P 112170-24-8P
112170-25-5P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation and esterification and antimicrobial activity of)
112170-20-4 CAPLUS
Leucine, N. N-(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl) sulfonyl) - (9CI)
(CA INDEX NAMS)

RN 112170-22-6 CAPLUS
CN Valine, N-{(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl}- (9CI)

(CA INDEX NAME)

112170-24-9 CAPLUS
Alanine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]- (9CI)(CA INDEX RAMS)

112170-25-9 CAPLUS L-Alanine, N-(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl)-(SCI) (CA IMDEX NAME)

Absolute stereochemistry.

112170-17-9P 112170-21-5P 112170-23-7P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation and esterification and peptide coupling and antimicrobial activity of)
112170-17-9 CAPLUS
L-Phenylelanine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl](9CI) (CA INDEX NAME)

Absolute stereochemistry.

112170-21-5 CAPLUS L-Leucine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]-(9C1) (CA INDEX NAME)

Absolute stereochemistry.

112170-23-7 CAPLUS L-Valine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Absolute stereochemistry.

112170-10-2 CAPLUS Leucine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]-, methyl ester (9C1) (CA INDEX NAME)

112170-11-3 CAPLUS
L-Leucine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]-,
methyl ester (9C1) (CA INDEX NAME)

Absolute stereochemistry.

112170-12-4 CAPLUS
Valine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]-, methyleater (9C1) (CA INDEX NAME)

112170-13-5 CAPLUS L-Valine, N-{(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl}-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

112170-14-6 CAPLUS
Alanine, N-[(1,2,3,4-tetrshydro-2,3-dioxo-6-quinoxalinyl)sulfonyl]-,
methyl ester (9C1) (CA INDEX NAME)

112170-15-7 CAPLUS L-Alanine, N-{[1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxalinyl)sulfonyl}-, methyl ester (9CI) (CA INDEX NAME)

112170-16-8P
RL: SPM (Synthetic preparation); PRSP (Preparation)
(preparation and peptide coupling and antimicrobial activity of)
112170-16-8 CAPUS

112170-16-8 CAPUS L-Tyrosine, N-[(1,2,3,4-tetrahydro-2,3-dioxo-6-quinoxaliny1)sulfony1]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

ACCESSION NUMBER:
ACCESSION NUMBER:
DOCUMENT NUMBER:
INVENTOR(S):
PATENT ASSIGNEE(S):
DOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
DATENT ASSIGNEE(S):
DOCUMENT TYPE:
DOCUM

DOCUMENT TYPE: LANGUAGE: FAMILY ACC, NUM, COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

| DE 3533791 | A1 | 19870326 | DE | 1985-3533791 | | 19850921 |
|------------------------|--------|-------------|----|--------------|---|----------|
| EP 216299 | A1 | 19870401 | EP | 1986-112816 | | 19860917 |
| R: CH, DE, FR, | IT, LI | | | | | |
| JP 62072678 | A2 | 19870403 | JP | 1986-219885 | | 19860919 |
| ES 2003353 | A6 | 19881101 | RS | 1986-2056 | | 19860919 |
| PRIORITY APPLN. INFO.: | | | DE | 1985-3533791 | A | 19850921 |
| OTHER SOURCE(S): | CASREA | CT 106:1964 | 59 | | | |
| | | | | | | |

IT

26773-25-1 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-ethyl- (8CI, 9CI) (CA INDEX NAME)

108229-84-1 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, 2-propenyl ester (9CI) (CA INDEX NAME)

108229-85-2 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, 2-ethoxyethyl ester (9CI) (CA INDEX NAME)

108229-86-3 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, 2-phenylethyl ester (9CI) (CA INDEX NAME)

108229-87-4 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, (4-methylphenyl)methyl ester (9C1) (CA INDEX NAME)

108229-88-5 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, (2-chlorophenyl)methyl ester (9C1) (CA INDEX NAME)

26773-32-0 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-chloroethyl)- (8CI, 9CI) (CA INDEX RAMB)

26921-20-0 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-2-propenyl- (9CI) (CA INDEX NAME)

108229-81-8 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, phenylmethyl ester (9CI) (CA INDEX NAME)

108229-83-0 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichlore-, butyl ester (9CI) (CA INDEX NAME)

108229-89-6 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, [4-{1,1-dimethylethyl)phenyl]methyl ester (9CI) (CA INDEX NAME)

108229-90-9 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, 2-chloroethyl ester (9CI) (CA INDEX NAME)

108229-92-1 CAPLUS 6-Quinoxalinecarboxamide, N-butyl-2,3-dichloro- (9CI) (CA INDEX NAME)

108229-94-3 CAPLUS 6-Quinoxalinecarboxamids, 2,3-dichloro-N-(phenylmethyl)- (9CI) (CA INDEX NAME)

108229-95-4 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-phenylethyl)- (9CI) (CA INDEX

108229-96-5 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(1-phenylethyl)- (9CI) (CA INDEX NAME)

108230-01-9 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, 1-phenylethyl ester (9CI) (CA INDEX NAMS)

108230-02-0 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, 2-furanylmethyl ester (9CI) (CA INDEX NAME)

108230-03-1 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-furanylmethyl)- (9CI) (CA INDEX NAME)

$$\overset{\circ}{\underset{\text{C1}}{\longrightarrow}} \text{CH}_2 - \text{NH} - \overset{\circ}{\text{C}} \overset{\cdot}{\underset{\text{N}}{\longrightarrow}} \overset{\cdot}{\underset{\text{C1}}{\longrightarrow}} \overset{\cdot}{\underset{\text{N}}{\longrightarrow}} \overset{\cdot}{\underset{\text{C1}}{\longrightarrow}}$$

Benzo[a]phenazine derivs. I (R1 = H, halo, Me, OH, alkoxy; R2 = CO2R5, COURGSR7; R3, R4 = H, alkyl; R5 = H, alkyl, cycloalkyl, Ph, PhcH2; R6, R7 = H, alkyl; NR6R7 = pyrrolidino, piperidino; n = 2, 3) are prepared as antitumor egents. A solution of eater II (R1 = OMe, R2 = CO2Me) in C6H6 was treated with H2NCH2CH2NNe2 and the mixture refluxed for 2 h to give I (R1 = OMe, R2 = CO2Me, R3 = R4 = Me, n = 2) (III). At 50 mg/kg/day i.p. for 5 days in mice transplanted with P3R8 leukenia cells. III increased survival time 5.88-fold, vs. 1.76-fold (maximum) for 5-FU at 25 mg/kg/day.
106224-71-99 106224-72-09 106224-78-69
106224-81-19 106224-87-79 106224-78-69
106224-81-19 106224-87-79 106224-78-69
(Reactant) or resgent)
(Reactant or resgent)
(Reactant or resgent)
(Preparation and amidation of)
106224-71-9 CAPUS
Benzo[a]phenazine-6,9-dicarboxylic acid, 10-ethoxy-5-hydroxy-, diethyl ester (9CI) (CA NUDEX NAME)

106224-72-0 CAPLUS Benzo[a]phenzin-6,9-dicarboxylic acid, 10-ethoxy-5-hydroxy-, 9-butyl 6-ethyl ester (9C1) (CA INDEX RAME)

106224-74-2 CAPLUS Benzo[a]phenazine-6,9-dicarboxylic acid, 5-hydroxy-10-propoxy-, diethyl eater (9CI) (CA INDEX NAME)

108258-55-5 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, (4-chlorophenyl)methyl ester (9C1) (CA INDEX NAME)

108258-56-6 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dichloro-, (4-methoxyphenyl)methyl ester (9C1) (CA INDEX NAME)

L13 ANSWER 105 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1987:50148 CAPLUS
DOCUMENT NUMBER: 206:50248 CAPLUS
106:50248 CAPLUS

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND DATE | APPLICATION NO. | DATE |
|------------------------|-------------------|---------------------|----------|
| | | | |
| EP 196910 | A2 19861008 | EP 1986-302395 | 19860401 |
| EP 196910 | A3 19870902 | | |
| EP 196910 | B1 19910102 | | |
| R: AT, BE, CH, | DE, FR, GB, IT, L | I, LU, NL, SE | |
| US 4686292 | A 19870811 | US 1986-838153 | 19860310 |
| CA 1248106 | A1 19890103 | CA 1986-504378 | 19860318 |
| JP 62000072 | A2 19870106 | JP 1986-64410 | 19860320 |
| JP 05013149 | B4 19930219 | | |
| ZA 8602183 | A 19861126 | ZA 1986-2183 | 19860324 |
| ES 553500 | A1 19870616 | ES 1986-553500 | 19860326 |
| AT 59642 | E 19910115 | AT 1986-302395 | 19860401 |
| PRIORITY APPLN. INFO.: | | JP 1985-65099 A | 19850329 |
| | | EP 1986-302395 A | 19860401 |
| OTHER SOURCE(S): | CASREACT 106:5024 | 8: MARPAT 106:50248 | |
| GI | | · | |

106224-75-3 CAPLUS
Benzo[a]phenazine-6,9-dicarboxylic acid, 5-hydroxy-10-propoxy-, 6-ethyl
9-propyl eater (9CI) (CA INDEX NAME)

106224-76-4 CAPLUS
Benzo[a]phenazine-6,9-dicarboxylic acid, 5-hydroxy-10-propoxy-, 9-butyl
6-ethyl ester (9C1) (CA INDEX NAME)

106224-78-6 CAPLUS Benzo(a)phenazine-6,9-dicarboxylic acid, 10-butoxy-5-hydroxy-, 9-butyl 6-ethyl ester (9CI) (CA INDEX NAME)

106224-61-1 CAPLUS
Benzo[a]phenazina-6,9-dicarboxylic acid, 5-hydroxy-, diethyl ester (9CI)

(CA INDEX NAME)

106224-87-7 CAPLUS Benzo[a]phenazine-6-carboxylic acid, 9-[(ethylamino)carbonyl]-5-hydroxy-10-methoxy-. ethyl ester (9CI) (CA INDEX NAME)

106224-90-2 CAPLUS
Benzo[a]phenazine-6,9-dicarboxylic acid, 5-hydroxy-10-metho
9-(2,2-dimethylpropyl) 6-ethyl ester (9CI) (CA INDEX NAME)

106224-68-4P 106224-69-5P 106224-91-3P 106224-94-91-3P 106224-94-6P 106224-96-6P 106225-00-7P 106225-01-P 106225-01-P 106225-01-P 106225-01-P 106225-01-P 106225-01-P 106225-12-P 106225-12-P 106225-12-P 106225-12-P 106225-12-P 106225-12-P 106225-12-P 106225-12-P 106225-12-P 106225-20-1P 106225-21-P 106225-22-3P 106225-20-P 106225-60-P 106225

106224-69-5 CAPLUS
Benzo[a]phenazine-9-carboxylic acid, 6-{{[a-(dimethylamino)ethyl]amino]carbonyli-5-hydroxyn-3-methylbutyl ester (SCI) (CA INDEX NOME)

106224-91-3 CAPLUS
Benzo[a]phenazine-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]carbonyl)-5-hydroxy-, ethyl ester (9CI) (CA INDEX NAME)

106224-94-6 CAPLUS
Benzo[a]phenazine-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]carbonyl]-5-hydroxy-10-methoxy-, 2,2-dimethylpropyl ester (9CI) (CA INDEX RAMS)

106224-96-8 CAPLUS Benzo[a]phenazine-6,9-dicarboxamide, N6-{2-(dimethylamino)ethyl}-N9-ethyl-5-hydroxy-10-methoxy- (9C1) (CA INDEX NAME)

106225-00-7 CAPLUS
Benzo[a]phenzalne-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]carboxyl]-10-ethoxy-5-hydroxy-, ethyl ester (9C1) (CA INDEX NAMS)

106225-01-6 CAPLUS
Benzo[a]phenazine-9-carboxylic acid, 6-{[[2-(dimethylamino)ethyl]amino]carbonyl]-10-ethoxy-5-hydroxy-, butyl ester (9CI) (CA INDEX NAMS)

106235-03-0 CADLUS

Benzo[a]phenazine-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]car

bonyl]-5-hydroxy-10-propoxy-, ethyl ester (9CI) (CA INDEX NAME)

106225-04-1 CAPLUS
Benzo[a]phenazine-3-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]carbonyl]-5-hydroxyne-0-propoxy-, propyl ester (SCI) (CA INDEX NAME)

106225-05-2 CAPLUS
Benzo[a]phenzolarie-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]carbonyl]-5-hydroxy-10-propoxy-, butyl ester (9CI) (CA INDEX RAME)

106225-07-4 CAPLUS
Benzo[a]phenazine-9-carboxylic acid, 10-butoxy-6-{{[2-(dimethylamino)ethyl]amino]carbonyl]-5-hydroxy-, butyl ester (9CI) (CA INDEX NAME)

106225-11-0 CAPLUS Benzo(a|phenazine-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]car bonyl)-5-hydroxy-10-methoxy-, sthyl seter [9CI] (CA INDEX NAME)

06225-12-1 CAPLUS enzo[a]phenazine-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]car onyl]-5-hydroxy-10-methoxy-, propyl ester (9CI) (CA INDEX NAME)

106225-13-2 CAPLUS Benzo(a|phenazina-9-carboxylic acid, 6-([[2-(dimethylamino)ethyl]amino)car bonyli-5-indydroxy-10-methoxy-, 1-methylathyl ester (9CI) (CA INDEX NAME)

106225-14-3 CAPLUS Benzo[a]phenazine-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]car bonyl)-5-hydroxy-10-methoxy-, 2-methylpropyl ester (9Cl) (CA INDEX NAME)

Benzo[a]phenazine-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]carbonyl]-5-hydroxy-10-methoxy-, octyl ester (9CI) (CA INDEX NAME)

106225-21-2 CAPLUS
Benzo[a]phenezine-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]carbonyli-5-hydroxyn-0-methoxy-, decyl ester (9CI) (CA INDEX RAME)

106225-22-3 CAPLUS

Benzo[a]phenzine-9-carboxylic acid, 6-{[[2-(dimethylamino)ethyl]amino]carbonyl]-5-hydroxy-10-methoxy-, pentadecyl ester (9CI) (CA INDEX NAME)

106225-23-4 CAPLUS
Benzo(a)phonaxin-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]carboxyl-1-5-hydroxyn-0-methoxy-, phenylmethyl ester (9CI) (CA INDEX NAME)

106225-15-4 CAPLUS
Benzo[a]phenszine-9-carboxylic acid, 6-{[{2-(dimethylamino)ethyl]amino]carbonyl|-5-hydroxy-10-methoxy-, pentyl ester (9CI) (CA IMDEX NAME)

106225-17-6 CAPLUS
Benzo[e]phenazine-9-carboxylic acid, 6-[[[2-(dimethylamino)ethyl]amino]car
bonyl[-5-hydroxy-10-methoxy-, hexyl ester (9CI) (CA INDEX NAME)

106225-19-8 CAPLUS
Benzo[a]phenarine-9-carboxylic acid, 6-[{[2-(dimethylamino)ethyl]amino]carbonyl]-5-hydroxy-10-methoxy-, heptyl ester (9CI) (CA INDEX NAME)

106225-20-1 CAPLUS

NH-CH2-CH2-NMe2

106225-25-6 CAPLUS
Benzo[a]phenazine-6,9-dicarboxamide, N6-[2-(dimethylamino)ethyl]-5-hydroxy-10-methoxy-N9-propyl- (9CI) (CA INDEX NAME)

106225-26-7 CAPLUS Benzo[a]phenas/ine-6,9-dicarboxamide, N9-butyl-N6-[2-(dimethylamino)ethyl]-5-hydroxy-10-methoxy- (9CI) (CA INDEX NAMS)

L13 ANSWER 106 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1987:19959 CAPLUS
DOCUMENT NUMBER: 106:19959
LITTLE: 106:19959
LITTLE: 106:19959
LITTLE: 106:19959
LITTLE: 106:19959
LITTLE: 106:19959
LITTLE: 206:19959
LITTLE: 206:19959
LITTLE: 206:19959
Moninger, Pritz: Schleefer, Ludwig

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE DE 3443305 EP 183142

| EP 183142 | A3 | 19890339 | | | |
|------------------------|----------|-----------|-----------------|---|----------|
| R: CH, DE, FR | , GB, IT | , LI | | | |
| IN 164645 | A | 19890429 | IN 1985-CA828 | | 19851121 |
| US 4707545 | A | 19871117 | US 1985-801751 | | 19851126 |
| JP 61130374 | A2 | 19860618 | JP 1985-265176 | | 19851127 |
| BR 8505929 | A | 19860819 | BR 1985-5929 | | 19851127 |
| CA 1253858 | A1 | 19890509 | CA 1985-496274 | | 19851127 |
| PRIORITY APPLN. INFO.: | | | DE 1984-3443305 | A | 19841128 |
| OTHER SOURCE(S): | MARPAT | 106:19959 | | | |
| GT | | | | | |

Li salts of anionic dyes are prepared by neutralizing an aqueous solution or suspension of an acidic dye with a Ca salt forming a low solubility Ca dye salt, which is treated with Li2SO4 or LiHSO4 to form the corresponding Li salt. Thus, a Na salt of an acidic monoare dye was reacted with CaCl2 to form a Ca salt containing filter cake, which was treated with a solution of LiSSO4 forming I, which was storage-stable at 50° for several weeks. The solution was spray-dried to form an electrolyte-containing fine dye nowder.

Li2SO4 forming I, which was storage-stable at 50° for several weeks. The solution was spray-dried to form an electrolyte-containing powder.

104601-66-3
RL: PROC (Proces)
(cation exchange of, with calcium chloride)
104601-66-3 CAPLUS
2-Anthracenssulfonic acid, 1-amino-4-[[4-{[[(2,3-dichloro-6-quinoxaliny]) carbonyl] methylamino|methyl]-3-sulfophenyl]manno]-9,10-dihydro-9,10-dioxo-, disodium salt (9CI) (CA INDEX NAME)

ΙT 106046-41-7

1986:610400 CAPLUS 105:210400 Storage-stable dye solutions Wolff, Joachim; Wolf, Karlheinz; Marschner, Nerner Bayer A.-G., Fed. Rep. Ger. Eur. Pat. Appl., 28 pp. CODEN: EPXXDW ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: Patent

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|-----|---------------------|--------|----------|-------------------|----------|
| | | | | | |
| | EP 167952 | A2 | 19860115 | RP 1985-108117 | 19850629 |
| | EP 167952 | A3 | 19890322 | | |
| | EP 167952 | B1 | 19910724 | | |
| | R: CH, DE, FR, | GB, LI | | | |
| | DE 3504964 | A1 | 19860123 | DE 1985-3504964 | 19850213 |
| | US 4685933 | A | 19870811 | US 1985-748267 | 19850624 |
| | JP 61036369 | A2 | 19860221 | JP 1985-149387 | 19850709 |
| | JP 05023306 | B4 | 19930402 | | |
| PRI | ORITY APPLN. INFO.: | | | DE 1984-3425813 A | 19840713 |
| | | | | | |

JP 05023306 84 19930402

PRIORITY APPIN. INFO:: DE 1984-3425813 A 19840713

AB Concentrated aqueous solns. of organic dyes and water-solubilizing agents contain

cyanamides N.tplbond.CNN2 or RC(:NH)NN2 (I; R * NHC.tplbond.N, NHCONN2, NHCONNH**). These aqueous solns. may contain an anionic reactive dye 7-35, a solubility-increasing water-miscible organic compound and/or hydrotrope and/or dispersing agent 0-30, N.tplbond.CNN2 or I 0.1-15, an inorg. salt 0-10, and a buffer 0-54. Thus, to 997.5 g aqueous suspension containing a sulfonated (phenylazolnaphthalene dye with a chlorodifluoropyrimidine group 11, inorg. salt mixture 1, and H3B03 0.54, 2.5 g dicyandiamide was added with stirring. A storage-stable aqueous solution was obtained showing no dye hydrolysis after 4 wk at 40°. This solution dyed rayon in a red-yellow tone.

In 10499-16-2

RN: USSS (Uses)

(concentrated aqueous solns. of, stabilizers against hydrolysis for)

RN 10499-68-2 CAPLUS

2-Anthracenesul fonic acid, 1-amino-4-[[4-[[(2,3-dichloro-6-quinoxalinyl)carbonyl]methylaminolmethyl]sulfophenyl|aminol-9, 10-dihydro-9, 10-dioxo-, monolithium monosodium selt (SCI) (CA INDEX NAME)

RL: PROC (Process)
(cation exchange of, with lithium oxalate)
16644-41-7 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[[4-[[(2,3-dichloro-6-quinoxaliny])carbonyl]methylamino]methyl]-3-aulfophenyllamino]-9,10-dihydro-9,10-dioxo-, calcium salt (9CI) (CA INDEX NAME)

106027-72-99
RL: IMF (Industrial manufacture); PREP (Preparation)
(preparation of, as storage-stable spray-dryable reactive dye composition)
106027-72-9 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[4-[[([2,3-dichloro-6-quinoxallnylcarbonyl]methylemino]methyl]-3-sulfophenyl]amino]-9,10-dihydro-9,10-dioxo-, dilithium salt (9CI) (CA INDEX NAME)

L13 ANSWER 107 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN

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D1- SO3H

● Li

● Na

L13 ANSWER 108 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DCCUMENT NUMBER:
105:210360
105:210360
Aqueous reactive dye solutions
Molff, Joachim; Molf, Karlheins; Seipt, Guenter
Bayer A.-Q., Fed. Rep. Ger.
GOURCE:
GOURCE:
CODEN: GONXBX
DOCUMENT TYPE:
Patent
LAWHIADE:
GONTAN

DOCUMENT TYPE: LANGUAGE: German PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DATE KIND APPLICATION NO. DATE DE 3424145 Al 19860109 DE 1984-3424145 19840630
PRIORITY APPLN. INFO::

B Aqueous solns. of reactive dyes containing water-solubilizing groups are
stabilized by addition of 0.1-25 R1(CR3R4)mC.tplbond.CCR5R6R2 [I; R1. R2
R4. R6 - H, C1-6 alkyl; R3. R5 - OH, (CCM2CR5/INDR) R7 - H, Ne; m - 0. 1;
n - 1-10). These solns. also contain a buffer 0-5, the reactive dye DE 1984-3424145 DE 1984-3424145

10-35, an inorg. salt 0-10, and a solubility-increasing water-miscible organic compound and/or hydrotrops and/or dispersing agent 0-30%. Thus, an aqueous solution was formulated containing a sulfated anthraquinone dye with a dichloroquinoxaline reactive group 20, e-caprolactam 10, MedNCCNSNBe 10, and a mixture of inorg, salts (NaCl, LiCl, Na2SO4, and LizSO4) 31. This solution (200 g) was mixed with 50 g urea, and 550 g water, giving solution A; filtrn. 15 s after addition of 200 g 20% NaCO3 showed dye. If to 800 g of solution A, 4 g I (m = 1, R1 = R2 = iso-Bu, R3 = R5 = Me, R4 = R6 = ON) in a mixture of ethylene glycol and polyethylene glycol alkylphenyl ether was added, followed by 200 g 20% NaCO3 solution, no signification

ipitation
was observed
88103-23-5D, lithium and sodium selts
RL: USSS (Uses)
(stabilizers for aqueous solns. of, alkynediols as)
88103-23-5 CAPUS
2-Anthracensulfonic acid, 1-amino-4-{(4-[{(2,3-dichloro-6-quinoxalinyl)carbonyl]methylamino]methyl]sulfophenyl]amino]-9,10-dioxo- (9CI) (CA INDEX EMMS)

PAGE 1-A

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D1-SO3H

ACCESSION NUMBER DOCUMENT NUMBER: TITLE:

AUTHOR(S): CORPORATE SOURCE:

SOURCE:

ANSWER 109 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
SSSION NUMBER:

1984:S10474 CAPLUS

1912:30474

LS:

Perivatives of 1,3-benzodioxole, 52. Preparation and reactions of 1,3-dioxolo[4,5-b] phenazines

Dallacker, Franz, Wagner, Alfred

Abt. Chem. Med., Tech. Hochach. Aachen, achen, p-5100, Fed. Rep. Ger.

RCE:

CE:

Lossier State State

$$\begin{array}{c} R \\ \\ \\ \\ \\ \\ \\ \end{array}$$

Naphthophenaxinones I (R = OMe, Me, NHAc, H, F, Cl, Br, CO2Et, CF3) were obtained by reaction of 6-substituted 1-cyclohexyl-2,3-dimethylquinoxalinium perchlorates with 2,3-dichloro-1,4-naphthoquinone [117-80-6]. The influence of R on general properties, lightfastness and on UV-visible and IR-NNR spectra is discussed. 67815-39.8

RU: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with dichloronaphthoquinone, ring formation in) 87815-93-8 CAPLUS

Cuinoxalinium 1-cyclohexyl-6-(sthoxycarbonyl)-2,3-dimethyl-, perchlorate

Quinoxalinium, 1-cyclohexyl-6-(ethoxycarbonyl)-2,3-dimethyl-, perchlorate (SCI) (CA IMDEX NAME)

CRN 87815-92-7 CMF C19 H25 N2 O2

2

CRN 14797-73-0 CMF C1 O4

L13 ANSWER 111 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1984:35784 CAPLUS
DOCUMENT NUMBER: 100:35784
TITLE: Concentrated liquid compositions of cold-dyeing

DOCUMENT TYPE:

Journal

German CASREACT 101:230474

$$\begin{array}{c} R^7 \\ R^5 \\ R^5 \\ R^6 \\ R^7 \\ N \\ N \\ R^2 \\ R^2 \end{array}$$

The 1,3-dioxolo[4,5-b]phenaxines I (R, R7 = H, Me, CO2Me; R1R2 = H2, COCHO; R3 = H, Me, CO2Me, R4 = H, CO2Me; R5 = H, CO2RE, Me, DMe; R6 = H, OVER; S8 = CO2Me; R6 = H, OVER; S8 = CO2Me; R5 = CO2Me; R6 = H, OVER; S8 = CO2Me; R6 = H, OVER; R6 = H, OV

L13 ANSWER 110 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1984:87233 CAPLUS DOCUMENT NUMBER: 100:87233

TITLE:

AUTHOR(S): CORPORATE SOURCE: SOURCE:

100:87233
Ring closure of quinonlymethane dyes and merocyanine analogs. Part 7. Synthesis and properties of 6-chloro-8-cyclohexyl-11-R-5,8-dihydronaphtho[1,2-bl]phenazin-5-ones Schelz, D.; Rottler, N. Inst. Farbenchem., Univ. Basel, Basel, CH-4056, Switz. Dyes and Pigments (1984), 5(1), 37-47 CODEN: DYPIDX; ISSN: 0143-7208
JOURNAL GRAND COMMENT OF THE PROPERTY OF

DOCUMENT TYPE:

LANGUAGE:

GI

| | fiber-reactive dyes |
|-------------------------|--|
| INVENTOR(S): | Hoquet, Robert G.; Kalz, Dietmar; Thomas, Thomas J.; |
| | Whetsell, Henry T.; Wolff, Joachim; Nonn, Konrad;
Wolf, Karlheinz |
| PATENT ASSIGNEE(S): | Bayer AG. , Fed. Rep. Ger.; Mobay Chemical Corp. |
| SOURCE: | Eur. Pat. Appl., 34 pp. |
| | CODEN: EPXXDW |
| DOCUMENT TYPE: | Patent |
| LANGUAGE: | German |
| FAMILY ACC. NUM. COUNT: | 1 |
| DATENT INFORMATION: | |

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|-----------|------------------|----------|
| | | | | |
| EP 92119 | A2 | 19831026 | EP 1983-103418 | 19830408 |
| EP 92119 | A3 | 19841107 | | |
| R: CH, DE, FR, | GB, LI | | | |
| US 4435181 | λ | 19840306 | US 1982-370426 | 19820421 |
| JP 58187460 | A2 | 19831101 | JP 1983-65768 | 19830415 |
| BR 6302071 | A | 19831227 | BR 1983-2071 | 19830420 |
| CA 1205253 | A1 | 19860603 | CA 1983-443149 | 19831213 |
| PRIORITY APPLN. INFO.: | | | US 1982-370426 A | 19820421 |
| OTHER SOURCE(S): | MARPAT | 100:35784 | | |

Storage-stable, aqueous cold-dyeing reactive dye compns. are prepared which contain 10-50 weights dye(s) with a fiber-reactive haloheterocyclic group and particle size <100 µ, sufficient enionic dispersant or polymeric N-vinyl lactam dispersant or prevent agglomeration or settling out of dye particles, and sufficient electrolyte to inhibit hydrolysis of the reactive group during temperature cycles ranging from 20° to 50°. A typical composition, stable for 3 wk during temperature cycles of 16 h at 20° and 8 h at 50°, contained dye I [78246-64-7] 31.5, lignosulfonate dispersant 3.0, NaCl 15.0, KM2PO4 0.2, KZHPO4 0.2, and HZO 50.03.

and RJO 50.0.
78246-64-7
RL: USEN (Uses)
(reactive dye, concentrated aqueous compns. containing, storage-stable)
78246-64-7
RJUS
2-Anthracenesul[onic acid, 1-amino-4-[[4-[[(2,3-dichloro-6-quinoxaliny])carbonyl]methylamino]methyllsul[ophenyl]metho]-9,10-dinydro3,10-dioxo, disodium selt (9C1) (CA INDEX RAMES)

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PAGE 2-A

D1- 503H

●2 Na

L13 ANSWER 112 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
INVENTOR(S):
INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:
Bayer A.-O., Fed. Rep. Ger.
SOURCE:
COURS. EPEXEN
LANGUAGE:
PATENT TYPE:
LANGUAGE:
PATENT HOPOPRATION:
PATENT HOPOPRATION:
COURS. EPEXEN
PATENT HOPOPRATION:
PATENT HOPOPRATION: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: APPLICATION NO. DATE

PATENT NO. KIND DATE

EP 87703 A1 19830907
EP 87703 B1 19850410
R: CH, DE, FR, OB, IT, LI
DE 3207534 A1 19830906
DE 3215933 A1 19831103
PRIORITY APPLN. INFO.: 19830221 EP 1983-101616 DE 1982-3207534 DE 1982-3215933 DE 1982-3207534 DE 1982-3215933 OTHER SOURCE(S): MARPAT 100:8505

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D1- 503H

87748-64-99
RL: PREP (Preparation)
(atorage-stable solution of, manufacture of)
87748-64-9 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[[4-[[[12,3-dichloro-6-quinoxalinyl]carbonyl]methylamino]methyl]mulfophenyl]mnino]-9,10-dihydro-9,10-dioxo-, dilithium salt (9CI) (CA INDEX NAMES)

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PAGE 2-A

D1- SO3H

L13 ANSWER 113 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1984:8427 CAPLUS
DOCUMENT NUMBER: 100:8427
TITLE: Preparations of water-soluble organic dyes
NONINTOR(S): Wolff, Joachim; Wolf, Karlheinz; Hoernle, Reinhold
Bayer A. O., Fed. Rep. Ger.
Ger. Offen., 23 pp.
CODEN: GMXXBX
DOCUMENT TYPE: PATENT LANGUAGE: PAPLLY ACC. NUM. COUNT: 1
FAMILY ACC. NUM. COUNT: 1 FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

Dyes containing an NH2 or C1-4-elkylemino group and 1-5 sulfo groups, and having salt content ≤8%, are treated with a chloro- and/or fluoro-substituted heterocyclic compound in aqueous medium in the presence of

acid-binding agent and, optionally, a solubilizer, and the resultant liquid is optionally buffered or dried and milled. The compns. are used to prepare dyeathen and printing pastes. For example, disactization of 0.3 mol. 2.5.1-HZN MINDCH2|CL0015CO3H, coupling with 0.3 mol 1.8.3.6-BXRH (RD)CL0014 (SO3H), salting out with 15 weight NIHAHCO3, weshing, and drying gave a dye [1] [70817-83-2] containing 34 malt. A soln of I in 700 mL H2O containing 3.25 mol c-caprolectam was treated with 0.3 mol 5-chloro-2.4.8-trifluoropyrimidine [697-83-6] in the presence of 0.32 mol LICCO3 and finally 0.13 mol CaO, and adjusted to pH 7 with 0.6 weight phosphate buffer to give a storage-stable solution containing 19 weight* II

.

88103-23-59
RL: PREP (Preparation)
(manufacture of, as powder composition with improved water solubility)
88103-23-5 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[[4-[[(2,3-dichloro-6-quinoxalinyl)carbonyl]methylamino]methyl]sulfophenyl]mmino]-9,10-dihydro-9,10-dioxo- (9CI) (CA INDEX NAMS)

PAGE 1-A

DATE 19830908 19830907 19850904 DE 3207533 EP 87705 EP 87705 DE 1982-3207533 EP 1983-101619 19820303 19830221 R: CH, DE, FR, GB, LI JP 58162667 A2 19830927 PRIORITY APPLN, INFO.:

Prepns. containing water-soluble anionic or cationic dyes and di-C1-4-alkyl sulfones are stable in storage. Thus, a mixture of reactive dye I [88112-50-9] 16.5, di-Me sulfone (II) [67-71-0] 10, phosphate buffer (for pH 7) 0.5, and H2O 73 parts was exirred and filtered to give a solution Storage of this solution for 4 wk at 50° resulted in only 44 hydrolysis of the dye, compared with 254 when II was replaced by tetramethylene sulfone.

78246-64-7 RL: USES (Uses)

RL: USES (Uses)
(aqueous solns. of, containing di-Me sulfone, storage-stable)
7246-64-7 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[[4-{[[(2,3-dichloro-6-quinoxaliny]) earbonyl] methylamino] methyl] sulfophenyl] amino]-9,10-dinydro-9,10-dioxo-, dieodium selt (9CI) (CA INDEX NAME)

PAGE 1-A

D1-503H

●2 Na

L13 ANSWER 114 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1983:596640 CAPLUS
DOCUMENT NUMBER: 99:196640
TITLE: 09:196640 CAPLUS
TITLE: NVENTOR(S): Wolfe, Joschine; Wolfe, Karlheinz; Ditzer, Reiner; Woernle, Reinhold Bayer A.-O., Fed. Rep. Ger.
Ger. Offen., 19 pp.
CODENS OMXEX
LANGUAGE: PATENT TYPE: PATENT LANGUAGE: OCCUMENT TYPE: CODENS OMXEX
DATENT HYPORATION: 9
TEATLY ACC. NUM. COUNT: 2
PATENT INFORMATION:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE 19830908 19830907 19850410 A1 DE 3207534 DE 1982-3207534 EP 1983-101616 DE 3207534 A1 19830906
EP 87703 A1 19830907
EP 87703 B1 19850910
R: CH, DE, FR, OB, IT, LI
JF 58162666 A2 19830927
JF 021008429 B4 19900302
ER 8301017 A 19831122
PRIORITY APPLM. INPO.: JP 1983-31079 19830228 BR 1983-1017 DE 1982-3207534 DE 1982-3215933 19830302 A 19820303 A 19820429 GI

AB Dyes of formula (MDIS)nQNHR (Q = dye residue; M = H, NN4, Li, Na, K; n = 1-5; R = H, C1-4 alkyl), with salt content 58 weights, are treated in aqueous media (optionally containing a solubilizing agent) with a halo heterocyclic compound in the presence of an acid acceptor and then optionally buffered to give a concentrated solution of reactive dye (MIOIS) nONRRI (R1 = fiber-reactive heterocyclic group; M1 = NN4, Li, Na, K; n and Q as defined above). For example, diszotization of 0.3 mol 4,3-H2N(HOS)5(CMSNIMAC [96-78-6], coupling with 0.3 mol 1-(4-sulfophenyl)-5-pyrazolone-3-carboxylic acid [118-47-8], deacetylation, dissoln, of the presscake by LiON in 450 mL H2O, treatment with 0.33 mol 2,3-dichloro-6-quinoxalinecarbonyl chloride [1919-43-3]

OTHER SOURCE(S):

CASREACT 99:196628

The title compds. (I; R = cyclohexyl, Ne; R1 = H2N. MeO, Me, AcNH, H, halogen. RCO2C, F3C. MeSO2, O2N), useful as precursors for naphthophenaxinone dyes were synthesized starting with 2-nitrohalobenzenes. The preferred method of condensing the diamine intermediate (II) with 2,3-butanedione [431-03-8] in mixts. of NOAc and HCLO4 was not successful whenever R1 was a strongly electron-withdrawing substituent. But in those cases R1 stabilized the corresponding enamines III, which could be obtained very easily.
97815-39-8P
RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and NOAC spectrum of)
87815-33-6 CAPIUS
Ouinoxalinium, 1-cyclohexyl-6-(ethoxycarbonyl)-2,3-dimethyl-, perchlorate IT

Ouinoxalinium, 1-cyclohexyl-6-(ethoxycarbonyl)-2,3-dimethyl-, perchlorate (8C1) (CA INDEX NAME)

CM 1

CRN 87815-92-7 CMF C19 H25 N2 O2

CRN 14797-73-0 CMF C1 04

while adding Li2CO3 to neutralize HCl, filtration, and buffering to pN 7 with 0.6% phosphate gave a storage-stable liquid crystalline preparation containing 22 weight% [87730-51-6].

If 87748-64-9 RL: USES (Uses)
(dye, manufacture of concentrated storage-stable solution of)
RN 87748-64-9 CAPLUS
CN 2-Anthracencesulfonic acid, 1-amino-4-[[4-([[(2,3-dichloro-6-quinoxalinyl)carbonyl]metholylamino]methyl]sulfophonyllamino]-9,10-dihydro-9,10-dioxo-, dilithium salt (9CI) (CA INDEX NAME)

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D1- SO3H

●2 Li

L13 ANSWER 115 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
1983:596628 CAPLUS
99:196628
Synthesis of 1-aryl- and 1-alkyl-2,3dimethylquinoxalinium perchlorates. Part 3.
Synthesis of 1,2,3-trimethyl-6-X- and
1-cyclohexyl-2,3-dimethyl-6-X- and
1-cyclohexyl-2,3-dimethyl-6-X- and
SOURCE:
COMPORATE SOURCS:
Schelz, D.; Rotzler, N.
Inst. Farbenchem., Univ. Basel, Basel, CH-4056, Switz.
Dyes and Pigments (1983), 4(4), 305-20
CODEN: DYPIDX; ISSN: 0143-7208
JOURNEL
LANGUAGE:
German

LI3 ANSWER 116 OP 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1983:181108 CAPLUS
COrrection of: 1983:55519
DOCUMENT NUMBER: 98:181108

DOCUMENT NUMBER:

TITLE:

98:181108
Correction of: 98:55539
Crossconjugated cyanines and merocyanines, obtained from salts of 1-substituted 2,3-dimethylquinoxalines. Part 2. Oxidative transformation of color bases Schelz, Dieter
Inst. Farbenchem., Univ. Basel, Basel, 4056, Switz. Helvetica Chimica Acta (1982), 65(5), 1607-16
CODEN: HCACAV; ISSN: 0018-019X
Journal
German

AUTHOR(S): CORPORATE SOURCE: SOURCE:

German

DOCUMENT TYPE: LANGUAGE: GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Products (I; R = Ph, 4-ClC6H4, Me; R1 = NO2, CF3, Br, H, OMe, etc.) of the readily oxidized II (X = N; R and R1 as defined) and II (X = N; R and R1 as defined) are sensitive to solvolysis, especially when R and R1 are electron-withdrawing substituents. In some cases, I could be identified as the oxidation products of III. The oxidation of III (R = Me, R1 = benzo.

as the oxidation products of III. The oxidation of III (R = Me, NI = benzo, A = 0.04 [52736-76-2] by alkaline K3Fe(CN) 6 leading to IV [84268-37-1] was compared with the voltammetric oxidation of V [68797-83-1] and related to the capto, dative-stabilized radicals proposed by N.O. Viehe et al. (1979). IH-NMR spectra of I were discussed with regard to 8-2 isomers. 84.25 p. (2010) [1979] [1970]

CM 1

Double bond geometry as shown.

CRN 14797-73-0 CMF C1 04

L13 ANSWER 117 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1983:55539 CAPLUS
DOCUMENT NUMBER: 58:55539 CAPLUS
S01:55539 CAPLUS
S01:55539 CAPLUS
S01:55539 CAPLUS
S01:55539 CAPLUS
Crosscoriugated cyanines and merocyanines, obtained from saits of 1-substituted 2,3-dimethylquinoxalines. Part 2. Oxidative transformation of color bases
Schelz, Dieter
CORPORATE SOURCE: Inst. Farbenchem., Univ., Basel, 4056, Switz.
Helvetica Chimica Acta (1982), 65(5), 1607-16
CODEN: HCACAV; ISSN: 0018-019X
JOURNAL LANGUAGE: German
OTHER SOURCE(S): CASRRACT 98:55539

OTHER SOURCE(S):

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

5,6-benzo.

A = ClO4 [52736-76-2] by alkaline K3Fe(CN)6 leading to IV [84268-37-1] was compared with the voltammetric oxidation of V [68797-83-1] and related to the capto, dative-stabilized radicals proposed by H. G. Viche et al. (1979). IH-NMR spectra of I were discussed with regard to E-Z isomers.

84267-32-3P (Properties); SPN (Synthetic preparation); PRSP (Preparation)
 (preparation and NNR spectrum of)
84267-32-3 CAPLUS
Quinoxalinium, 2,2'-(1,2-ethenediyl)bis[6-(ethoxycarbonyl)-3-methyl-1-

using the Ames test. DX1 (I) [81485-18-9] was potentially mutagenic in Salmonella typhimurium TA 100 and 98 with and without the S-9 mixture WO 25 [81485-17-8] And WO 20 [81485-16-7], being atructurally related to I, did not show any genetic change in the strains used. The antibiotic activity of these chems. was also tested using gram-neg, and gram-pos. bacteria. I had more killing effect in gram-pos. bacteria than WO 25 and WO 20.

WO 20.
81485-16-7
RL: ADV (Adverse effect, including toxicity); BIOL (Biological study)
(mutagencity and toxicity of)
81485-16-7 CAPLUS
D-Phenylelanine, N-[[3-[[[18,2R]-2-hydroxy-1-methyl-2-phenylethylmethylamino]cerbonyl]-2-methyl-1,4-dioxido-6quinoxalinyl]carbonyl]- (9CI) (CA INDEX NAME)

ANSWER 119 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

CAPLUS COPYRIGHT 2006 ACS on STN
1982:87009 CAPLUS
96:87009
Cross-conjugated cyanines and merocyanines, obtained
from selze of 1-substituted 2,3-dimethylquinoxalines.
Part 1. Isolation of the dye bases from spontaneous
transformation or oxidation of the reactants with
copper(II) acetate or silver oxide
Schelz, Dieter
Inst. Parbenchen., Univ. Basel, Basel, CH-4056, Switz.
Helvetics Chimica Acta (1981), 64(8), 2665-80
CODEN: HCACAV; ISSN: 0018-019X
JOURNAL

AUTHOR (S):

CORPORATE SOURCE: SOURCE:

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI

Journal German CASREACT 96:87009

Quaternary salts I (R=Me, Ph, p-ClC6H4; R1=H, electron acceptor or donor; R2=Me, Ph; X=CH, NI, in some cases in the presence of the corresponding II, undergo spontaneous conversion to III (all groups as defined for I) when dissolved in DNSO or DMP. Yields are 24-479. Higher

phenyl-, (E)-, diperchlorate (9CI) (CA INDEX NAME)

CRN 84267-31-2 CMF C38 H34 N4 O4

Double bond geometry as shown.

CRN 14797-73-0 CMF Cl O4

AUTHOR (S):

L13 ANSWER 118 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE:
Microbial mutagenicity and toxicity of newly
synthesized heterocyclic N-oxides
AUTHOR(S):
Al-Mossevi, M. A. J.; Salema, A. A.; Salema, M.; Anani,

CORPORATE SOURCE:

A. Kuwait Inst. Sci. Res., Safat, Kuwait Environment International (1981), 5(3), 141-4 CODSN: ENVIOV: ISSN: 0160-4120 Journal English

DOCUMENT TYPE: LANGUAGE:

Newly synthesized heterocyclic N-oxides were tested for their mutagenicity

yields (up to 66%) are obtained by oxidation of I, II, or I-II mixts. with Cu(OAc)2 or Ag2O. Visible and IH-MRR spectra data for the dyes are given, and their structural relationship to S. Huenig's (1980) two-step redox systems is discussed. 68763-65-1.

RL: RCT (Reactant); RACT (Reactant or reagent) (oxidative dimerization of) 68765-65-1 CAPLUS (Quinoxalinium, 6-(ethoxycarbonyl)-2,3-dimethyl-1-phenyl-, perchlorate (SCI) (CA INDEX NAME)

IT

CM 1

CRN 68765-64-0 CMF C19 H19 N2 O2

CRN 14797-73-0 CMF C1 O4

L13 ANSMER 120 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1981:497719 CAPLUS
DOCUMENT NUMBER: 55:97129
The Synthesis of "stretched-out" analogs of lumazine,
6.7-disethyllumazine and 3-amino-5, 6.7, 6-tetrshydro6.7-disethyl-4-pteridinone
Schneller, Stewart N., Christ, William J.
Dep. Chem., Univ. South Floride, Tampa, FL, 31620, USA
JOURGE: JOURNE JOURNE

DOCUMENT TYPE: LANGUAGE: GI

Treating 2.4.5-(H2N)3C6H2CO2Et with glyoxal and McCOCOMe gave I (R = H, Me), which were treated with urea to give pyrazinoquinarolinedione II.

III was similarly prepared
78795-09-27 78795-109-7
RD: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation and cyclocondensation of, with urea)
78795-09-2 CAPLUS
6-Quinoxalinecarboxylic acid, 7-amino-, ethyl ester (9CI) (CA INDEX NAME)

78795-10-5 CAPLUS 6-Quinoxalinecarboxylic acid, 7-amino-2,3-dimethyl-, ethyl ester (9CI) (CA INDEX NAME)

L13 ANSWER 121 OF 101 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1981:463663 CAPLUS
DOCUMENT NUMBER: 95:63663 CAPLUS
1NVENTOR(S): 2,3-Dichloroquinoxaline-6-carboxamide derivatives
Gleinig, Harald; Lahrs, Juergen; Jovcic, Dorde;
Schubert, Klaus; Goom, Walter; Goesling, Claus
BAFENT ASSIGNEE(S): 8ayer A.-G., Fed. Rep. Ger.
COUNTRY TYPE: Patent
DOCUMENT TYPE: Patent

DOCUMENT TYPE: Patent German 1

LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

APPLICATION NO. PATENT NO. DATE DE 2942364 DE 2942364 PRIORITY APPLN. INFO.: DR 1979-2942364 19791019 A1 C2 19810423 19861120 DE 1979-2942364 A 19791019

Title compds. (I; Z = organic radical; R = H, Cl-4 alkyl; n = 1-4) are prepared in high yield by treating Z (BMR)n in aqueous medium at 5-20° and pH 3.5-5 with molten 2,3-dichloroquinoxaline-6-carboxyl chloride [1919-43-3] at 110-180° which is introduced beneath the surface of the aqueous phase via a spray nox1e. The method is especially useful for preparing I in

DOCUMENT NUMBER:

Tis a dye residue, e.g., azo, anthraquinone, or phthalocyanine. The preparation of II [78246-64-7] and several other fiber-reactive dyes is described.
78181-07-47 78246-64-7P
RL: IMF (Industrial manufacture); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses) (dye, manufacture of)
78181-07-4 CAPLUS
Cuprate(3-), [3-[4-[[5-([[(2.3-dichloro-6-quinoxalinyl]carbonyl]methylami nolesthyl]-2-sulfophenyl]azo]-2-hydroxy-3-methylphenyllazo]-4-hydroxy-2,7-naphthalenedisulfonato(5-)]-, trisodium (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

78246-64-7 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[{4-{[[(2,3-dichloro-6-quinoxaliny])earbony] methylamino]eathyl]sulfophenyl]amino]-9,10-dihydro-9,10-dioxo-, disodium salt (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

D1-503H

●2 Na

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. APPLICATION NO. EP 1980-302411 PATENT NO.

1 23785
EP 23785
E KIND DATE DATE 1981011 EP 1980-302411 19810429 19850403 , GB, IT, NL, SE 19841108 AU 1980-59547 DE, FR, 19790717 A1 A1 A1 BABCA1 B2 A2 B4 B1 19810806 ZA 1980-3928 IL 1980-60506 CA 1980-356027 HU 1980-1762 19800630 19800706 19800711 19800715 19810806 19810624 19861231 19930316 19830928 19850729 19810118 19910311 DK 1980-3068 19800716 19910311 19910819 19810127 19810701 19860116 19860930 19810414 19940223 19850415 BR 1980-4413 ES 1980-493431 CS 1980-5044 SU 1980-2951003 JP 1980-96960 19800716 19800716 19800716 19800716 19800717 AT 1980-302411 US 1981-334384 US 1986-939694 DK 1989-1684 19800717 19850415 19870407 19890207 19890407 19940321 19890407 19911111 19920330 19811224 19861209 19890407 DK 1989-1685 19890407 AU 1979-9617 AU 1980-3093 US 1980-164933 EP 1980-302411 A 19790717 A 19800411 A2 19800701 A 19800717 A 19810112 A3 19811224 AU 1981-7201 US 1981-334384 OTHER SOURCE(S): CASREACT 95:25134

L13 ANSWER 122 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1981:425134 CAPLUS

The title compds. I (X = optionally substituted OC6H4O, OC6H4S, SC6H4S; R = H, optionally substituted alkyl, scyl; R1 = H. optionally substituted alkyl; RR1 = H. optionally seterified CO2H, substituted Me; R3 = H. halogen, cyano, carbanoyl, optionally seterified CO2H, substituted Me; R3 = H. halogen, cyano, thiocyano, optionally substituted NH2, aliphatic, OH, SH, CO2H, or CONH2; m, n = 0, 1; p = 0-2) were prepared Thus, 2,6-dichloroquinoxaline was treated with 4-HOC6H4CCHMeCO2HM to give 70% II. At 1 kg/ha preemergence II gave 100% control of ryegrass and Japaness millet.
78104-80-0P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
78104-80-0 CAPLUS
6-Ouinoxalinecarboxylic acid, 3-[4-(2-ethoxy-1-methyl-2-oxoethoxy)phenoxy]-, ethyl ester (9CI) (CA INDEX NAME)

$$\bigcup_{B \in O-C}^{0} \bigcup_{N = 0}^{Me} \bigcup_{\substack{O-CH-C-OE}}^{0} \bigcup_{N = 0}^{Me} \bigcup_{N = 0}^{0} \bigcup_{N = 0}^{N} \bigcup_$$

L13 ANSWER 123 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1981:141191 CAPLUS
DOCUMENT NUMBER: 94:141191
TITLE: Diseaze copper complex dyes
INVENTOR(S): Jacger, Horst
DATENIT ASSIGNES(S): Seyer, Horst
SOURCE: Ger Offen, 11 pp.
COLMENT TYPE: CANGUAGE: GERANDER
LANGUAGE: NUMBER COLDEN: GERANDER

COLMENT TYPE: GERANDER

COLMENT T

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DATE 19810122 19810218 19830921 APPLICATION NO. KIND DE 2925210 EP 23955 EP 23955 DE 1979-2925210 EP 1980-103234 R: CH, DE, FR, GB
JP 56005858 A2
PRIORITY APPLN. INFO.: 19810121 JP 1980-82214 DE 1979-2925210 19800619 A 19790622

PAGE 1-B

L13 ANSWER 124 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1981:92561 CAPLUS DOCUMENT NUMBER: 94:92561 TITLE: Studies of the state of

AUTHOR(S): CORPORATE SOURCE: SOURCE:

94:92561
Studies on sulfensnlides. V. Anodic exidation of 4'-substituted 2-nitrobenzenesulfensnlides at a reticulated vitreous carbon electrode Sayo, Hiroteru; Mori, Koichi; Michida, Takashi Fac. Pharm. Sci., Kobe-Gakuin Univ., Kobe, 673, Japan Chemical: Pharmaceutical Bulletin (1980), 28(12), 3707-10
CODEN: CPBTAL; ISSN: 0009-2363
JOURNAL BROLIA

DOCUMENT TYPE:

DOCUMENT TYPE: Journal
LANGUAGE:

AB Constant current electrolysis of 4'-R-2-nitrobenzenesulfenanilides (I; R = OMe, Me, Cl. COZE) was carried out in MeCN containing 0.1M ethyltributylammonium trifluoromethanesulfonate, 1% trifluorometic acid, and 1% trifluorometic analyside at a reticulated vireous C (RVC) electrode. The quantity of electricity to be fed into the electrolytic coll was determined from the anodic potential vs. time curves. The yields of 2,7-di-R-phenazines (R = OMe, Me, Cl, and COZEC) were 56, 24, 42, and 33%, resp. The RVC anode was useful for electrolysis of I because the considerable yields of phenazines were obtained within several minutes without using an expensive potenticetat.

IT 72848-45-49
RL: PREP (Preparation)
(preparation of, by electrochem. oxidation of ethoxycarbonylnitrobenzenesulfena nilide on glassy carbon in acetonitrile)
RN 72848-45-4 CAPUS

N 72848-45-4 CAPUS

L13 ANSWER 125 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1980:93992 CAPLUS 92:93992 TITLE: Studies - Studies -

Studies on sulfenamides. IV. Oxidation of 4'-, 3'-and 2'-substituted 2-nitrobenzenesulfenanilides with

Disazo copper complex dyes (I; R = H, Cl-4 slkyl; Rl = H, Me, Et, OMe, OEt; Z = 1- or 2-hydroxynaphthalenesulfonic acid derivative residue) are manufactured by coupling dissocited 3,4-HBN(HOJS)CEHICHENERRE (R defined as above, R2 = seyl component) with a coupling component 4,2-R1(HBN)CEHIORINGRIQRI (R1 defined as above, R3 = Me, Et), dissociting the aminosso intermediate, coupling with the hydroxynaphthalenesulfonic acid derivative, deskylatively copperizing and removing R3, and condensing with 2,3-dichloro-6-quinoxalinecerbonyl chloride [19:49-43-]. Thus, I (R = R1 = Me, Z = Z1, azo bond in 2-position, OH in 1-position) [77000-78-3] was prepared by this method.
77000-78-35 CAPLUS (R: LST) [1-(15-(((2,3-dichloro-6-quinoxalinyl)carbonyl]methylami nojmethyl]-3-sul[cphenyl]asol-3-hydroxy-5-methylphenyl]asol-4-hydroxy-2,7-naphthalenediaulfonato(5-)]-, trihydrogen (9CI) (CA INDEX NAME)

PAGE 1-A

AUTHOR(S):

AUTHOR(S):

Sayo, Hiroteru; Mori, Koichi; Michida, Takashi
CORPORATE SOURCE:

Fac. Pharma Sci., Kobe-Gakuin Univ., Kobe, 673, Japan
Chemical & Pharmaceutical Bulletin (1979), 27(10),
2316-20

CODEN: CPBTAL; ISSN: 0009-2363

DOCUMENT TYPE:

JOURNAL
LANGUAGE:

English
AB The oxidation of 2-02NC6H4SNNC6H4R [R = 4-Br [I], 4-COZE [II], 4-COMe [III],
4-OEE [IV], 4-NO2 [V], 4-SO2NH2 [VI], 3-Me [VII], 2-OMe [VIII], and 2-Me
(IX)] by PbO2 was carried out in MeCN containing 1% CF3CO2H and 1% (CF3CO)2O.
The oxidation of I-V and VII gave 2, 7-disubstituted phenazines, whereas that
of VI, VIII and IX did not. AcNNC6H4NO3-2 was obtained in all cases and
(2-02NC6H48)2 was obtained from II-IV, VI, and VIII. The oxidation of IX
gave a small amount of 2'-methyl-N-[(2-nitrophenyl)thio]-pbenroquinometimie, while that of VIII gave a mixture of 2'-methoxy-N-[(2nitrophenyl)thio]-o- and -p-benzoquinometimes.

1 72848-45-4 CAPLUS
CN 2,7-Phenazinedicarboxylic acid, diethyl ester [9CI] (CA INDEX NAME)

L13 ANSWER 126 OP 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
171TLE:
1NVENTOR(S):
Okamoto, Shosuke; Kikumoto, Ryoji; Tamao, Yoshikuni;
Okubo, Kazuo; Tezuka, Toru; Tonomura, Shinji;
Hijikata, Akiko
Mitsubishi Chemical Industries Co., Ltd., Japan

PATENT ASSIGNEE (S): SOURCE: Ger. Offen., 147 pp. CODEN: GNXXBX Patent

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|-------------|------|----------|-----------------|----------|
| | | | | |
| DE 2801478 | A1 | 19780720 | DE 1978-2801478 | 19780113 |
| DE 2801478 | C2 | 19910131 | | |
| US 4066773 | A | 19780103 | US 1977-760745 | 19770119 |
| US 4073913 | A | 19780214 | US 1977-760668 | 19770119 |
| US 4093712 | A | 19780606 | US 1977-760672 | 19770119 |
| US 4097472 | A | 19780627 | US 1977-760676 | 19770119 |
| US 4101653 | A | 19780718 | US 1977-760929 | 19770119 |
| US 4097591 | A | 19780627 | US 1977-776195 | 19770310 |
| JP 54003037 | A2 | 19790111 | JP 1977-66508 | 19770606 |
| JP 60010028 | 84 | 19850314 | | |
| US 4125604 | A | 19781114 | US 1977-804334 | 19770607 |
| US 4131673 | A | 19781226 | US 1977-804368 | 19770607 |
| US 4140681 | A | 19790220 | US 1977-804331 | 19770607 |

| IL 53685 | A1 | 19851231 | | | 19771223 |
|--------------------------|--------------|---|--|-----|--|
| AU 7832289
AU 522320 | A1 | 19790719 | AU 1978-32289 | | 19780109 |
| ZA 7800123 | B2
A
A | 19790829 | ZA 1978-123
PI 1978-73 | | 19780109 |
| FI 7800073 | A | 19780720 | PI 1978-73 | | 19780110 |
| PI 72316
PI 72316 | В | 19870130 | | | |
| ES 466706 | A2 | 19781016 | RS 1978-466706 | | 19780110 |
| NL 7800448 | A | 19870130
19870511
19780721
19910801
19920102
19780720
19871207
19880317
19820628
19830228
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19870710
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19820515 | ES 1978-466706
NL 1978-448 | | 19780113 |
| NL 187746 | В | 19910801 | | | |
| NL 187746
SE 7800512 | C | 19920102 | SE 1978-512 | | 19780117 |
| SE 452624 | B | 19871207 | 00 1370 011 | | |
| SE 452624 | ç | 19880317 | | | |
| HU 22709
HU 180265 | 0 | 19820628 | HU 1978-MI626 | | 19780117 |
| DK 7800263 | Ä | 19780720 | DK 1978-263 | | 19780118 |
| DK 150521 | B | 19870316 | | | |
| DK 150521 | c | 19871019 | | | |
| NO 7800191
NO 158681 | A | 19780720 | NO 1978-191 | | 19780118 |
| NO 158681 | č | 19861019 | | | |
| FR 2378004 | A2 | 19780818 | FR 1978-1368 | | 19780118 |
| FR 2378004
GB 1596971 | B2 | 19850913 | GB 1978-2063 | | 19780118 |
| PL 123267 | A
81 | 19821030 | PL 1978-204063 | | 19780118 |
| CH 633773 | Ä | 19821231 | CH 1978-519
CH 1978-4530 | | 19780118 |
| CH 648293 | A | 19850315 | CH 1978-4530 | | 19780118 |
| SU 1181539
BE 863092 | A3 | 19850923 | SU 1978-2566652 | | 19780118 |
| ES 466705 | A2 | 19790816 | CH 1978-4530
SU 1978-2566652
BE 1978-184463
BS 1978-466705
DD 1978-203302
AT 1978-399 | | 19780119 |
| DD 137352 | c | 19790829 | DD 1978-203302 | | 19780119 |
| AT 7800399 | A | 19820515 | AT 1978-399 | | 19780119 |
| AT 369356
CS 236757 | B2 | 19821227 | CS 1978-381 | | 19780119 |
| JP 62014548 | B4 | 19870402 | JP 1978-4529 | | 19780119 |
| JP 54100342 | A2 | 19790808 | | | |
| US 4173630
SU 938739 | A . | 19791106 | US 1978-902855
SU 1979-2776611 | | 19780504
19790618 |
| AT 8003284 | Ä | 19820515 | AT 1980-3284 | | 19800623 |
| AT 369357 | В | 19821227 | | | |
| AT 8003285
AT 369358 | A | 19820515 | AT 1980-3285 | | 19800623 |
| CS 236772 | B2 | 19850515 | CS 1981-2011 | | 19810319 |
| CS 236773 | B2 | 19850515 | CS 1981-2011
CS 1981-2012 | | 19810319 |
| FI 8402539 | A | 19840621 | PI 1984-2539 | | 19840621 |
| FI 74455
FI 74455 | B | 19871030
19880208 | | | |
| IORITY APPLN. INFO | | .,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,, | US 1977-760668 | A | 19770119 |
| | | | US 1977-760672 | A | 19770119 |
| | | | US 1977-760676
US 1977-760745 | A | 19770119
19770119 |
| | | | US 1977-760929 | | 19770119 |
| | | | | _ | 19770310 |
| | | | JP 1977-66508 | À | 19770606
19770607 |
| | | | US 1977-776195
JP 1977-66508
US 1977-804331
US 1977-804368
JP 1974-128774 | Α . | 19770607 |
| | | | JP 1974-128774 | Ä | 19741108 |
| | | | | A | 19741108 |
| | | | JP 1974-136695 | A | 19741129
19741129
19750225
19750226
19750305 |
| | | | JP 1974-136697
JP 1975-23268
JP 1975-23635 | Ä | 19750225 |
| | | | JP 1975-23635 | A | 19750226 |
| | | | JP 1975-26768
JP 1975-29357 | A | 19750305 |
| | | | OF 19/5-29357 | A | 19750311 |
| | | | | | |

| JP | 1975-29358 | A | 19750311 |
|----|-------------|-------|----------|
| US | 1975-62 | 239A3 | 19751014 |
| US | 1975-622390 | A3 | 19751014 |
| US | 1975-638985 | A2 | 19751209 |
| US | 1976-646522 | A | 19760105 |
| US | 1976-649219 | | 19760114 |
| US | 1976-653217 | A2 | 19760128 |
| US | 1976-656014 | A | 19760206 |
| US | 1976-656870 | A | 19760210 |
| US | 1976-669743 | A | 19760324 |
| US | 1976-671436 | A2 | 19760329 |
| υs | 1976-671568 | A2 | 19760329 |
| US | 1976-703704 | A2 | 19760708 |
| US | 1976-707536 | A2 | 19760722 |
| US | 1976-713486 | A2 | 19760811 |
| US | 1976-723474 | A | 19760914 |
| US | 1976-728051 | A | 19760930 |
| US | 1977-760677 | A2 | 19770119 |
| PI | 1978-73 | A | 19780110 |
| CH | 1978-519 | Ä | 19780118 |
| AT | 1978-399 | Ä | 19780119 |
| CS | 1978-381 | A3 | 19780119 |
| | | | |

OTHER SOURCE(S): MARPAT 90:152610

Absolute stereochemistry.

L13 ANSMER 127 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCRSSION NUMBER: 1979:18673 CAPLUS
DOCUMENT NUMBER: 90:38673
THILE: 90:38673
AUTHOR(S): 20:38673 CAPLUS
AUTHOR(S): 20:38673 CAPLUS
CORPORATE SOURCE: 30:48674 CAPLUS
DOCUMENT TYPE: 40:48674 CAPLUS COPEN HEACAV; ISSN: 0018-019X
JOURNAL DOCUMENT CAPLUS CAPL

OTHER SOURCE(S):

A general method for the preparation of the title compds. I (X = CH, R = Ph, Rl = NO2, 802Me, CN, CP3, CO2Ec, Cl. Br, Me, OMe; R = Me, 4-ClC6He, Rl = NO2, H, X = CH, R = Ph, Rl = H, X = N) involved the condensation of 4.2-R(H2N)C6H3NHR with MeCOCOMe and HclO4 in a mixed solvent containing excess St2O. I were converted into II by heating with Bt3N and Me2CO. The NOMR shifts of I were correlated with Hammet's constant op. I are useful

anites of I very correlated with Hammet's constant op. I are useful as dye precursors.
68765-65-19
KI: PRP (Properties); SPN (Synthetic preparation); PRSP (Preparation) (preparation and NRR of)
67655-65-1 CAPLUS
GNINOKALINIUM, 6-(ethoxycarbonyl)-2,3-dimethyl-1-phenyl-, perchlorate (SCI) (CA INDEX NAME)

CM 1

CRN 68765-64-0 CMP C19 H19 N2 O2

CRN 14797-73-0 CMF Cl O4

L13 ANSWER 128 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1978:120955 CAPLUS
BOCUMENT NUMBER: 88:120955 CAPLUS
SOME reactions of 2,3-diaminoindole derivatives.
Synthesis of indolo[2,3-b]quinoxalines
Kurilo, G. N.; Rostowa, N. 1.; Cherkasova, A. A.;
Grinev, A. N.
CORPORATE SOURCE: Vees. Nuchmo-Issled. Khim.-Farm. Inst. im.
Ordzhonikidze, Moseow, USSR
Khimiya Geterotsiklicheskikh Soedinenii (1977), (12),
1645-7
CODEN: KOSSAQ; ISSN: 0453-8234
JOURNEL LANGUAGE: Ruesian
OTHER SOURCE(S): CASREACT 88:120955

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): GI

Treatment of indoles I (R = H, O- and p-Me, p-Cl. p-CO2Et; R1 = H, Me) with Cu acetate in MeOH gave 40-64% II (R2 = MeO); II (R = H, p-Cl, R1 = H; R2 = piperidino) were obtained in 45-51% yield in the presence of piperidine. Heating II (R = H, O-Me, p-Cl, p-CO2Et; R1 = H; R2 = OMe) at 200-10 $^\circ$ (F0 $^\circ$ min gave 15-33 $^\circ$ III (R3 = H, 4-Me, 2-Cl, 2-CO2Et)

III (R3 = 2-Cl, 2-CO3Et) monoxides were obtained in 61-87% yield by oxidation of III by H2O2. 65880-42-4P
RU: SPM (Synthetic preparation); PREP (Preparation) (preparation of) 65880-42-4 CAPLUS 6H-Indolo[2,3-b]quinoxaline-2-carboxylic acid, 6-methyl-, athyl ester (SCI) (CA INDEX NAME)

L13 ANSWER 129 OF 161 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1977:553420 CAPLUS
87:153420 2.4+Disryl[1,3,4H]thiadiszines fueed to quinoxalines
HINDENTOR(8):
BILIOIT. Arthur John
du Pont de Nemoure, E. I., and Co., USA
U.S., 10 pp.
COORN: USXXAM
PAECHT
P

Patent English

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|-------|--------------|------------------|----------|
| | | | | |
| US 4025510 | A | 19770524 | US 1975-636792 | 19751201 |
| PRIORITY APPLN. INFO.: | | | US 1975-636792 A | 19751201 |
| GI For diagram(s), see | print | ed CA Issue. | | |

GI For diagram(e), eee printed CA Issue.

AB 2,4-Diaryl-13,4-thiadisarines (I, R = Ph, substituted Ph, 4-quinolinyl; RI = Ph, substituted Ph, 4-quinolinyl; RI = Ph, substituted Ph; A = quinoxaline, pyramine, pyrimidine, pyridine, pyrididine, pyri

| OF 32030730 | ~ | 13//0314 | OF 13/0-1303 | 80 | 19/01101 |
|------------------------|----|----------|--------------|-------|----------|
| GB 1540604 | A | 19790214 | GB 1976-4550 | 2 | 19761102 |
| GB 1540605 | A | 19790214 | GB 1977-2250 | 2 | 19761102 |
| FR 2330738 | A1 | 19770603 | FR 1976-3348 | 9 | 19761105 |
| FR 2330738 | B1 | 19800808 | | | |
| CH 624426 | A | 19810731 | CH 1977-1222 | 1 | 19771006 |
| PRIORITY APPLN. INFO.: | | | DE 1975-2549 | 570 A | 19751105 |
| | | | CH 1976-1382 | 1 A | 19761102 |
| | | | GB 1976-4550 | 2 A | 19761102 |

10220514

TR 1076 130500

RZNHCH2

TD F00F4770

Fiber-reactive azo dyes [I. R = 5-chloro-2,6-difluoro-4-pyrimidiny], 2,3-dichloro-6-quinoxaliny]; R1 = naphthelenesulfonic acid, pyrazole, acetoacetanilide, azo chromophore residue; Z = direct bond, CO] were prepared and used to dye and print cotton and wool fast yellow to blue shades. Thus, -1-anino-2-sulfo-3-(aninomethy)1-4-methoxybenzene [63353-60-6] was prepared, diazotized, coupled with 1,3,6-HOC10HS(SOH)2 [578-85-8], the resulting azo compound treated with 2,4,6-trifluoro-5-chloropyrimidine [697-83-6], and the reaction mixture salted to give I(R = 5-chloro-2,6-difluoro-4-pyrimidiny], R1 = 1,3,6,2-HO(ROS)2C10HR; Z = direct bond] [63353-68-4], dyeing cotton a fast yellowish red shade. IT

63355-43-7
RI: TEM (Technical or engineered material use); USES (Uses)
(dye, for cellulosic fibers, preparation of)
63355-43-7
CAPLUS
CUPTATE(3-), [3-[[4-[[3-[[(2,3-dichloro-6-quinoxalinyl)carbonyl]amino]met
hyl]-4-methoxy-2-eulophenyl[azo]-2-hydroxy-5-methylphenyl[azo]-4-hydroxy2,7-naphthalenedieulfonato(5-]]-, trihydrogen (9CI) (CA INDEX NAME)

PAGE 1-A

63811-22-3 CAPLUS
1H-[1,3,4]Thiadiazino[5,6-b]quinoxaline-7-carboxamide,
N,N-diethyl-1,3-diphenyl- (9CI) (CA INDEX NAME)

IT

26807-34-3 63810-79-7
RL: RCT (Reactant); RACT (Reactant or reagent)
(reaction of, with (thiobenzoyl)phenylhydrazine)
26807-34-3 CAPLUS
6-Quinoxalinecarboxamide, 2,3-dichloro-N,N-diethyl- (&CI, 9CI) (CA INDEX NAME)

63810-79-7 CAPLUS 6-Quinoxalinecarbo INDEX NAME) exalinecarboxamide, 2,3-dichloro-N-(3-methoxypropyl)- (9CI) (CA

87:69732
Piber-reactive azo dyes
Jaeger, Horat
Bayer A.-G., Fed. Rep. Ger.
Ger. Offen, 49 pp.
CODEN: GMXXEX
PALENT
German
1 INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------|------|----------|-----------------|----------|
| | | | | |
| DE 2549570 | A1 | 19770512 | DE 1975-2549570 | 19751105 |
| DE 2549570 | C2 | 19830519 | | |

63353-62-8

RL: TEM (Technical or engineered material use); USES (Uses)
(dye, for cotton, preparation of)
63353-62-8 CAPLUS

Benzenesulfonic acid, 6-{[1-(2-chloro-5-sulfophenyl)-4,5-dihydro-3-methyl-5-oxo-1H-pyrazol-4-yllazol-2-[[(2,3-dichloro-6-quinoxalinyl)carbonyl]amino]methyl]-3-methoxy- (9CI) (CA INDEX NAME)

L13 ANSWER 131 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1977:89889 CAPLUS
DOCUMENT NUMBER: 86:89889
TITLE: Azepino(4,5-b)quinoxalines
Hurmaus, Rudolf; Oriss, Gerhart; Grell, Molfgang;
Sauter, Robert; Reichl, Richard; Leitold, Matyas
FATENT ASSIGNEE(S): Thomae, Dr. Karl, G.m.b.H., Fed. Rep. Ger.
COCUMENT TYPE: COCUMENT ONXXBX

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. DATE 19750430 A 19750430 KIND DATE DE 2519258 PRIORITY APPLN. INFO.: A1 19761111 DE 1975-2519258 DE 1975-2519258

Tetrahydro-1H-azepino(4,5-b)quinoxalines (I; R = e.g., H, Me, Ph. PhCH2, Ac. Bz, CO2H, CH2CM2CO2H; Rl = Rl = H, OH, AcO, Eco2CO; RZn = e.g., H, s-Cl, 7-NO2, 8-Me, 8-CO2H, 8,9-Me2, 8-MeO), useful as appetite depresents and bactericides (no data), are prepared by various known methods, mostly involving reaction between an o-phenylenediamine and an exepinedione. The atepinedione can be obtained by cyclization of an iminodipropionic acid derivative Thus, reaction of PhCH2N(CH2, CH2CO2Me) with Na and Me38iCl Inde2C6H4 gives 1-benzyl-2,3,6,7-tetrahydro-4,5-big(trimentylsiloxy)-IH-azepine which is exidized with Br to the azepine-4,5-dione which then reacts with 1,2-(H2N)2C6H4 in AcOH to give after 4 hr at 100° 834 I.HCl (R = PhCH2, Rl = R2n = R3 = H).
6.1793-13-07
RL: SPN (Synchatic preparation); PREP (Preparation) (preparation of)
6.1793-152-0 CAPLUS
IH-Azepino (4,5-b)quinoxaline-8-carboxylic acid, 2,3,4,5-tetrahydro-3-(phenylmethyl)-, ethyl estar (9CI) (CA INDEX NAME)

L13 ANSWER 132 OF 181 CAPLUS COPYRIGHT 2006 ACS on ETN ACCESSION NUMBER: 1976:412370 CAPLUS DOCUMENT NUMBER: 5:12370
TITLE: Stable polymer factors

Stable polymer images by photopolymerization in a matrix

INVENTOR(S): PATENT ASSIGNEE(S): SOURCE:

matrix
Baumann, Niklaus
Ciba-Geigy A.-G., Switz.
Ger. Offen., 109 pp.
CODEN: GWXXBX
Patent

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

DATE PATENT NO. APPLICATION NO KIND DE 2525674 CH 594704 CH 604208 DE 1975-2525674 CH 1974-7956 CH 1974-7957 FR 1975-17754 19760102 19780131 19750609 19780831 FR 2274951 FR 2274951 CA 1077760 19760109 19831028 19750606 CA 1975-228849 BE 1975-157171 JP 1975-71385 19750609

-Quinoxalinecarboxamide, N,N-diethyl-2,3-diphenyl- (8CI, 9CI) (CA INDEX VAME)

37966-43-1 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-bis[(acetyloxy)methyl]-, ethyl ester [GCI] (CA INDEX NAME)

52996-75-5 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-bis(bromomethyl)-, ethyl ester (9CI) (CA INDEX NAME)

L13 ANSWER 133 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1975:423265 CAPLUS
39:22365
Rifect of 1-phenazinecarboxylic acid derivatives on
experimental tumore
AUTHOR(8):
Sidorik, O. A.: Shevchenko, I. N.
CORPORATE SOURCE:
Fixiologicheski Aktivnyo Veshchestva (1966-1992)
(1974), 6. 92-4
CODEN: FAVUAI; ISSN: 0533-1153

US 1976-743011 CH 1974-7956 CH 1974-7957 CH 1975-4843 US 1975-584444 19770823 US 4043819 PRIORITY APPLN. INFO.: ат

Photopolymerizable compns. capable of forming stable polymer relief images are composed of an ethylenically unsatd. photopolymerizable monomer, a chemical hardenable, nonlight-sensitive, swellable macromol. compound as binder, a hardening agent, and a quinoxaline derivative (1; R = H, Me; R1 = Me, Ph, NaOlSCSH4, p-MeOCSH4; R2 = H, SO3H), which together with the monomer or the binder forms a redox pair, as photoinitator. Thus, a gelatin-subbed cellulose triacetate support was overcoated with a solution containing calcium discrylate 26.07, acrylamids 3.89, galatin 7.58, glycerin 2.14, the ether of polyethylene glycol with N-(methylol) perfluorosityleulifonamide 0.107, B-(1,5-dimethyl-1-pyrazoly) lacrolein (hardener) 0.081, and I (R, R2 = H; R1 = NASOJCSH4) 1.03 g/s2, dried at 30°, contact exposed under a photog, step wedge (12 steps) for .apprx.30 sec to a 400°H high-pressure Hg lamp at 40 m, rinsed with water, colored with a cationic dyce rinsed with water, and dried to show 12 steps amaximum color d. was 2 cm and the absolute and cried to show 12 steps and 10 maximum color d. was 2 cm and 10 m and 10 m

CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dimethyl-, ethyl ester (6CI, 8CI, 9CI) (CA INDEX NAME) ÇN

RN 32388-08-2 CAPLUS CN 6-Quinoxalinecarboxamide, N,N-diethyl-2,3-dimethyl- (8CI, 9CI) (CA INDEX

DOCUMENT TYPE: Journal
LANGUAGE: Russian

GI For diagram(s), see printed CA Issue.

AB When injected s.c. at 200 mg/kg/day into rats for 12-15 days or i.p. at 50
mg/kg/day into mice for 8-12 days, 1-phenazinecarboxylic acid Na salt [1]
[134-02-1] or glycine N-1-phenazinecarboxyl acid Na salt [5]
53327-47-4] significantly inhibited the growth of Ehrlich
carcinome and erythromyelosis. The inhibitory effects of the prepns. on
lymphoma NK/Ly or sarcoma 45 were less pronounced and both compds. were
inactive sgainst Guerin carrinoms.

IT 53327-47-4
RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); TRU (Therapeutic use); BIOL (Biological study); USES
(Uses)
(neoplasm inhibitor)

(Dees) (neoplasm inhibitor)
55327-47-4 CAPUS
Olycine, N-(2-phenazinylcarbonyl)-, monosodium salt (9CI) (CA INDEX NAME)

● Na

L13 ANSWER 134 OF 181 CAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1975:156377 CAPLUS DOCUMENT NUMBER: 82:156377

TITLE: INVENTOR(S):

82:156377
Piperazinyl quinoxalines
Engelhard. Edward L.; Lumma. William C., Jr.; Saari,
Walfred S.
Merck and Co., Inc.
Ger. Offen., 36 pp.
CODEN: GWXEX
Patent
German PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

> DATE APPLICATION NO. DATE 19740711 19740625 19740626 19740627 DE 2433397 PI 7401939 DX 7403426 NO 7402351 SE 7408486 SE 417316 SE 417316 NI 7408705 AU 7470731 GB 1440722 ES 428107 PR 2236499 19750206 DE 1974-2433397 FI 1974-1939 DK 1974-3426 NO 1974-2351 SE 1974-8486 A1 A A A B C A A1 A1 A1 C A1 19750114 19750303 19750114 19750114 19810309 19810625 19750115 19760108 19760623 NL 1974-8705 AU 1974-70731 GB 1974-30176 ES 1974-428107 FR 1974-24114 DD 1974-179871 BE 1974-146519 ZA 1974-4466 CH 1974-9648 19740702 19740708 19740709 19740711 19740712 19740712 19740712 19761116 PR 2236499 DD 112127 BE 817608 19750207 19750320 19750113

JP 1974-79774 US 1973-379022 US 1974-465381 JP 50037791 19750408 19740713 PRIORITY APPLN. INFO.:

For diagram(e), see printed CA Issue.

Piperaxinylquinoxalines I (R = N, Me, COMMe, COCHARC, CH2CH2ON, Ac, COUSCM-2, COCYCHAP, CSCCHIZODER, e. llyl, CH2CHCHCI; R1 = N, Cl, Me, COZH, Ph, COZEC, SPh, Ac, NNCH2CH2ON, NH2, OSt; R2 = halo, NO2, OMe, CP3 etc. in S-6 positione) and some related compds. (50 compds.) vere prepared for use as antidepressants, appatite depressants, and analysiscs. Thus reaction of 2,3-dichioro-6-cyanoquinoxaline with N-formylpiperaxine and reduction over Pd-C gave I (R = R1 = N, R2 = 6-CN).

\$5668-57-2P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

\$5566-57-2 CAPLUS
6-Ouinoxalinecarboxamide, 3-chloro-N-(2-hydroxyethyl)-2-[4-(2-hydroxyethyl)-1-piperaxinyl]-, monohydrochloride (SCI) (CA INDEX NAME)

● HC1

26773-13-7 RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with piperazine derivs.) 26773-13-7 CAPLUS

xalinecarboxamide, 2,3-dichloro-N-(2-hydroxyethyl)- (8CI, 9CI) (CA

L13 ANSWER 135 OF 101 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1974:570266 CAPLUS
DOCUMENT NUMBER: 91:170266 Photopolymerization of ethylenically unsaturated compounds
INVENTOR(5): Baumann, Niklaus; Schlunke, Hans P.
CODUMENT ASSIGNEE(S): Ciba-Geigy A.-O.
COUNTY TYPE: COUNTY OF THE COU

FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO.

KIND DATE

APPLICATION NO.

PR 2221453
B1 19780908
CH 1972-17658
R 19721205
PRIORITY APPLN. INFO.:
CH 1973-18758
CH 1972-17658
A 19721205
CH 1973-18714
A 19731205
CH 1973-18714
CH 1972-17658
A 19721205
CH 1973-18714
A 19731207

AB Quinoxaline derive, and salts (.sim.150) were described which were useful with electron donors, such as Na p-toluenseulfinet (1) [824-79-3] and triphenylphosphine (603-35-0], as photoredox catalysts for the preparation ofsacrylaside-bearium discrylate copolyper [37281-67-7] (sep., on photogo, film supports) or a similar polymer upon exposure to radiation (200-450 mm). Thus, a solution of 1.4 M aqueous Ba diacrylate 180, 1.6 M aqueous acrylamide
60, 6\$ aqueous gelatin 30, and 0.25\$ aqueous FC 170 (wetting agent) 30 ml was mixed with 2 ml ECOM containing 10 mg 6.7-ethylenedioxy-2,3-bis(hydroxymethyl)quinoxaline (II) [52995-38-0] and 2 ml 0.016 M aqueous I, costed on gelatin-coated cellulose triacetate film, dried, irradiated with a Hg lamp through a photog, step-wedge for 30 sec, and developed with a dys to give discernible shades corresponding to the steps.

13 2328-09-3 37966-43-1 52996-73-5
RL: CAT (Catalyst use); USSS (Uses) (catalysts, contg electron donors, for photopolymn)
RN 3238-05-9 CAPLUS
CN 6-Quinoxalinecarboxylic acid, 2,3-diphenyl-, ethyl ester (8CI, 9CI) (CA INDEX NAME)

6-Quinoxalinecarboxylic acid, 2,3-dimethyl-, ethyl ester (6CI, 8CI, 9CI) (CA INDEX NAME)

32386-08-2 CAPLUS 6-Quinoxalinecarboxamide, N,N-diethyl-2,3-dimethyl- (8CI, 9CI) (CA INDEX NAME)

Quinoxalinecarboxamide, N,N-diethyl-2,3-diphenyl- (8CI, 9CI) (CA INDEX

43-1 CAPLUS
OXAlinecarboxylic acid, 2,3-bis[(acetyloxy)methyl]-, ethyl ester
(CA INDEX NAME) 6-Quinc (9CI)

52996-75-5 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-bis(bromomethyl)-, ethyl ester (9CI) (CA INDEX NAME)

AUTHOR (S) :

LI3 ANSWER 136 OF 161 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1374:146079 CAPLUS
BOILDON'S E0:146079 CAPLUS
FOILER: 80:146079
Nuclear magnetic resonance studies of heterocyclic bridged biphenyls
AUTROR(S): Hall D. Muriel; Hwang, Huaun-Yong; Bhanthumnavin,

Biravana Dep. Chem., Bedford Coll., London, UK CORPORATE SOURCE:

Journal of the Chemical Society, Perkin Transactions 2: Physical Organic Chemistry (1972-1999) (1973), (15), 2111-CODEN: JCPKBH; ISSN: 0300-9580

Journal English DOCUMENT TYPE:

NAGE: English
For diagram(s), see printed CA Issue.
Condensation reactions between 1,2-dismines and biphenyl-2,2'dicarboxaldehyde or 9,10-phenanthraquinone, and between
2,2'-disminobiphenyl and 1,2-diketones, gave polycyclic products with 5-,
6-, 7-, and 8-membered heterocyclic rings, the NRR spectra of which are
discussed. The 15H-dibenzo[c,e]benizimidazo[1,2-a]azepines I (R * H,
COSEC) are fluxional. The dibenzo[a,c]phenazines II showed large
downfield shifts (\$\delta\$.apprx.9.5) for some aromatic protons.
\$1448-1.745

IT

downfield shifts (8 .apprx.9.5) for some aromatic protons.
51448-17-4Petic preparation); PREP (Preparation)
(preparation of)
51448-37-4 CAPLUS
Dibenzo[a,c]phenazine-11-carboxylic acid, ethyl ester (9CI) (CA INDEX NAME)

L13 ANSWER 137 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1974:109835 CAPLUS
60:109835
Ac reactive dyes
Jacger, Horst
Bayer A.-G.
CODN: GRXXEX
DOCUMENT TYPE:

LARGE CAPLUS COPYRIGHT 2006 ACS on STN
1974:109835
CAPLUS CAPLUS
60:109835
CAPLUS

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| • | | | | | |
|---|-------------|------|----------|-----------------|----------|
| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
| | | | | | |
| | DE 2232541 | A1 | 19740117 | DE 1972-2232541 | 19720703 |
| | DE 2232541 | B2 | 19771027 | | |
| | IT 990815 | A | 19750710 | IT 1973-26026 | 19730628 |
| | BE 801661 | A1 | 19740102 | BE 1973-132901 | 19730629 |
| | JP 49052828 | A2 | 19740522 | JP 1973-73009 | 19730629 |
| | JP 55043025 | B4 | 19801104 | | |
| | CH 739539 | A4 | 19750530 | CH 1973-9539 | 19730629 |
| | CH 572546 | B | 19760213 | | |
| | CA 994330 | A1 | 19760803 | CA 1973-175278 | 19730629 |
| | CH 582739 | A | 19761215 | CH 1975-15476 | 19730629 |
| | NL 7309200 | A | 19740107 | NL 1973-9200 | 19730702 |
| | DD 107302 | c | 19740720 | DD 1973-171990 | 19730702 |
| | RS 416498 | A1 | 19760301 | ES 1973-416498 | 19730702 |
| | GB 1431322 | A | 19760407 | GB 1973-31420 | 19730702 |
| | GB 1431323 | A | 19760407 | GB 1975-39759 | 19730702 |
| | AT 320100 | В | 19750127 | AT 1973-5845 | 19730703 |
| | FR 2236905 | Äl | 19750207 | FR 1973-24415 | 19730703 |
| | | | | | |

| US | 4126609 | A | 19781121 | US | 1973-376184 | | 19730703 |
|----------|---------------|----|----------|----|--------------|-----|----------|
| AT | 7400179 | A | 19760515 | AT | 1974-179 | | 19740110 |
| AT | 334313 | В | 19760110 | | | | |
| JP | 52063488 | A2 | 19770525 | JP | 1976-11493 | | 19760206 |
| JP | 57029592 | B4 | 19820623 | | | | |
| US | 4049704 | A | 19770920 | US | 1976-656251 | | 19760209 |
| PRIORITY | APPLN. INFO.; | | | DE | 1972-2232541 | Α | 19720703 |
| | | | | AT | 1973-5845 | . A | 19730703 |
| | | | | US | 1973-376184 | A3 | 19730703 |
| | | | A | | | | |

Ano and diseato fiber-reactive dyes (I, R = 1 - (sulfophenyl) -4-pyrazolyl derive., aminohydroxyaulfonaphthyl derive., (sulfophenyl) -4-pyrazolyl derive., R1 = 2.6-diffuoro-5-chloro-4-pyrimidinyl; 2,6-dichloroquinoxaline-6-carbonyl) were prepared and were used to dye cotton fast greenish yellov to navy blue shades. Thue, 2,1-AcMICIORSONS in N2508 was treated with N-(hydroxymethyll)phthalimide at 15-20.deg. for 24 hr and the product treated with aqueeus NaOH at 170.deg. to give 2-amino-5-(aminosethyl)-1-naphthalenseulfonic acid (52084-84-1) which was diszotized and coupled with 1,8,3,5-MG(BRN)[1048 (503H) 2 to give an azo intermediate (III) treatment of II with 2,4,6-crifluoro-5-chloropyrimidine gave azo reactive due (III) [51366-30-4]. The other I were similarly prepared 52084-97-4P
RL: IMP (Industrial manufacture); PREP (Preparation) (preparation of)

(preparation of)

S2084-87-4 CAPLUS

1-Nephthe lensulfonic acid, 2-[[1-(2-chloro-5-sulfophenyl)-4,5-dihydro-3-sethyl-5-oxo-li-pyrasol-4-yl] axol-5-[[(2,3-dichloro-6-quinoxalinyl)-earbonyl]-danisolsethyl]- (SCI) (CA INDEX NAME)

LI3 ANSWER 138 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1973:418659 CAPLUS
DOCUMENT NUMBER: 79:18669 Reaction of sodium borohydride with heteroaromatic nitro compounds
AUTHOR(S): Reaction of sodium borohydride with heteroaromatic nitro compounds
AUTHOR(S): New York Compounds
ALTHOR(S): ARO, KOPPAK V.; Jackman, Dennis
J. Hills Miller Health Cent., Univ. Florida, Gainesville, Ft, USA
Journal of Heterocyclic Chemistry (1973), 10(2), 213-15
COUMENT TYPE: OUDIN: JNTCAD; ISSN: 0022-152X
JOURNAL
DOCUMENT TYPE: JUNEAU ACCESSION OF ACCES

40859-07-2P
RL: IMF (Industrial manufacture); PREP (Preparation)
(preparation of)
40859-07-2 CAPLUS
2H-Naphtho[1,2-d]triazole-5,9-disulfonic acid, 7-[[1-[[(4-amino-3,5-disulfophenyl]amino]carbonyl]-2-oxopropyl]azol-2-[4-[[[(2,3-dichloro-6-quinoxalinyl)carbonyl]methylamino]methyl]phenyl]- (9CI) (CA INDEX NAME)

L13 ANSWER 140 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1972:552222 CAPLUS
TITLE: Substituted quinoxalines for the inhibition of gastric acidity
INVENTOR(S): Bolhofer, William A.; Baldwin, John J.
PATENT ASSIGNER(S): Merck and Co., Inc.
U.S., S pp.
COUMENT TYPE: LANGUAGE: PAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

US 3691166 A 19720912 US 1969-886791 19691219

PRIORITY APPLIN. INPO.:
GI For diagram(s), see printed CA Issue.

8 Seven 2.3-dichloro([2-bydroxyethyl)carbamoyl]quinoxalines {I, when R = CONN(CH2)20H, R 1 = H, Cl, R2 = H, Cl, Me, R3 = H, Cl, se; when R1 = CONN(CH2)20H, R = R2 = H, R3 = Cl, Me, R3 = H, Cl, se; when R1 = prepared by refluxing the 2.3- and 3.4-diaminobenzoic acids with di-Et oxalate, converting the resulting 2.3-dihydroxy-50r 61-quinoxalinecarboxylic acids with PCLS-POCL3 to the 2.3-dichloroquioxalinecarboxyl chorides and treating these with NO(CH2)2NH2 act room temperature

IT 31901-98-79 37902-03-7P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
RN 37901-98-7 CAPLUS

of II resp. 5-Nitroisoquinoline was reduced to the 1,2,3,4-tetrahydro derivative in HOAc at 5° but yielded the 1,2-dihydro derivative in aqueous

MeOR.

8924-72-7
RL: RCT (Reactant); RACT (Reactant or reagent)
(reduction of, by sodium borohydride in acetic acid)
6924-72-7 CAPLUS
6-Quinoxalinecarboxylic acid, ethyl ester (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)

L13 ANSWER 139 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1973:73615 CAPLUS
1973:73615 C

DOCUMENT TYPE:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|-----------------------|------|----------|-----------------|------------|
| | | | | |
| DE 2113298 | A | 19720921 | DE 1971-2113298 | 19710319 |
| DE 2113298 | B2 | 19770512 | | |
| DE 2113298 | C3 | 19771229 | | |
| NL 7203510 | A | 19720921 | NL 1972-3510 | 19720316 |
| BE 780848 | Al | 19720918 | BE 1972-115221 | 19720317 |
| FR 2130422 | A5 | 19721103 | PR 1972-9516 | 19720317 |
| FR 2130422 | B1 | 19751024 | | |
| IT 953527 | A | 19730810 | IT 1972-22046 | 19720317 |
| AT 309619 | В | 19730827 | AT 1972-2310 | 19720317 |
| DD 102159 | č | 19731212 | DD 1972-161627 | 19720317 |
| AT 316479 | В | 19740710 | AT 1972-7973 | 19720317 |
| GB 1378244 | Ä | 19741227 | GB 1972-12626 | 19720317 |
| CA 998388 | A1 | 19761012 | CA 1972-137324 | 19720317 |
| CH 606341 | A | 19781031 | CH 1974-8948 | 19720317 |
| JP 54027020 | B4 | 19790907 | JP 1972-26718 | 19720317 |
| ES 400909 | Al | 19750116 | ES 1972-400909 | 19720318 |
| US 4118382 | A | 19781003 | US 1977-800573 | 19770525 |
| RIGRITY APPLN. INFO.: | • | 22,0200 | | 19710319 |
| RIORIII AFFEM: INFO:: | | | | 1 19720317 |

DE 1971-2113398 A 19710319

4'-Amino-1', 5'-disulfoacetoacetanilide 1972-238565 Al 19720317

4'-Amino-1', 5'-disulfoacetoacetanilide (T) [37615-83-1] was used as the coupling component to prepare 6 fiber-reactive are dyes [II, 0 = phenyl, naphthotriazolyl, or benzothiazolyl containing a fiber-reactive group) which dyed cotton light- and wetfast yellow to greenish yellow shades. Thus, 1,4,2,6-(H2M)268H2(SOSH)2 was condensed with diketene in aqueous NAOH to give 1. 2,5-H2M/ORN)C6H303H was disactized and coupled with I, the intermediate nitro azo compound reduced with Na2S, and the amino derivative condensed with 3,3-dichloroquinoxaline-6-carbonyl chloride to give azo dye III [37615-47-7], greenish yellow on cotton. In another typical example, 2-(methylaulfonyl)-6-methoxy-7-aminobenzothiazole was diazotized and coupled with I to give azo dye IV [37615-48-8], yellow on cotton.

6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-hydroxyethyl)-6-methyl- (9CI) (CA INDEX NAME)

37902-03-7 CAPLUS 6-Quinoxalinecarboxamide, 2,3,8-trichloro-N-(2-hydroxyethyl)- (9CI) (CA INDEX NAME)

ACCESSION NUMBER:

ACCESSION NUMBER:

DOCUMENT NUMBER:

TITLE:

Substituent effects on coupling constants in bicyclic heterocanatic compounds and the prediction of chemical shifts from coupling constants.

AUTHOR(S):

AUTHOR(S):

AUTHOR SOURCE:

SOURCE:

SOURCE:

SOURCE:

CONFORATE SOURCE:

SOURCE:

CONFORATE SOURCE:

CONFORM:

CONFORATE SOURCE:

CONFORM:

(substituent coupling consts. in, substituent chemical shift in relation to)
6924-72-7 CAPLUS
6-Quinoxalinecarboxylic acid, ethyl ester (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)

L13 ANSWER 142 OF 181 CAPLUS COPYRIGHT 2006 ACS on 6TN
ACCESSION NUMBER: 1972:503277 CAPLUS
TITLE: 7:103277
Pher reactive dyes
INVENTOR(8): Dehmel, Georgy Jaeger, Horet
Partent Assignes(8): 6er. Offen. 86 pp.
CODEN: GXXXXX
DOCUMENT TYPE: Patent GXXXXX

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---|---------------------|------|----------|-------------------|----------|
| | | | | | |
| | DE 2055967 | A | 19720518 | DR 1970-2055967 | 19701113 |
| | IT 951598 | A | 19730710 | IT 1971-30972 | 19711111 |
| | CH 549632 | A | 19740531 | CH 1973-4115 | 19711111 |
| | BE 775265 | A1 | 19720301 | BE 1971-110436 | 19711112 |
| | NL 7115598 | A | 19720516 | NL 1971-15598 | 19711112 |
| | FR 2114670 | A5 | 19720630 | FR 1971-40675 | 19711112 |
| | GB 1369856 | A | 19741009 | GB 1971-52667 | 19711112 |
| | JP 56003389 | 84 | 19810124 | JP 1971-89976 | 19711112 |
| ı | ORITY APPLN. INFO.: | | | DE 1970-2055967 A | 19701113 |
| | | | | | |

JP 56003389 B4 19810124 JP 1971-89976 19711112
ORITY APPLM. INFO.: DE 1970-2055967 A 19711112
FOUTEER H2O-soluble fiber-reactive azo, phthalocyanine, nitro, and anthraquinone dyes containing 502MH802(GH2)MMMEX groups (X = 2,3-dichloroquinoxalin-6-carbonyl or 5-chloro-2,6-difluoropyrimidin-4-yl, n = 3 or 4) were prepared and used to dye cellulose and wool wetfast shades. For example, p-H3NC6H8003MN502CH2CH2CH2CH2MHWe.fer. 1-(2-chloro-4-sulfophenyl)-3-methyl-5-pyrazolone was condensed with 2,3-dichloroquinoxaline-6-carbonyl chloride to give fiber-reactive dye I (35914-00-0).
159914-00-01 38997-34-6P 38097-35-7P
RE: HW (Industrial manufacture); PREP (Preparation)
(preparation of)
159934-00-0 CAPLUS
Benzenesulfonic acid, 3-chloro-4-[4-[[4-{[[3-[(2,3-dichloro-6-quinoxalinyl)carbonyl]methylamino]propyl]sulfonyl]amino]sulfonyl]phenyl]az
o)-4,5-dihydro-3-methyl-5-oxo-1H-pyrazol-1-yl]- (SCI) (CA INDEX NAME)

PAGE 2-A

D1-503-

38097-35-7 CAPLUS
Cuprate(1-), (C. (mainosulfony1)-C,C-bis[[[4-[[(2,3-dichloro-6-quinoxaliny1)carbony1]methylamino|buty1]sulfony1]amino|sulfony1]-29H,31H-phthalocyanine-C-sulfonato(3-)-N29,N30,N31,N32]-, hydrogen (9CI) (CA INDEX NAME)

PAGE 1-A

38097-34-6 CAPLUS
Cuprate(1-), [C-(aminosulfonyl)-C,C-bis[{[[4-[[(2,3-dichloro-6-quinoxalinyl)carbonyl]methylamino|butyl]sulfonyllamino|sulfonyl]-29H,31H-phthalocyanine-2-sulfonato(3-)-N29,N30,N31,N32]-, hydrogen [9CI] (CA

PAGE 1-A

PAGE 2-A

D1-503-

36153-46-7 CAPLUS
2-Maphthalenesulfonic acid, 7-(acetylamino)-3-[[3-[[[3-{[[2,3-dichloro-6-quinoxalinyl]oarbonyl]methylamino]propyl]sulfonyl]amino]sulfonyl]phenyl]az
o]-4-hydroxy- (9CI) (CA INDEX NAME)

PAGE 1-B

L13 ANSNER 143 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1972:461948 CAPLUS
DOCUMENT NUMBER: 77:61948
Synthesis and properties of some phenazine derivatives and their N-mono- and N,N-dioxides
AUTHOR(S): Return R. Kh.; Komyukhov, V. N.; Pushkareva, Z. V.;
Yarysheva, I. A.
CORPORATE SOURCE: Ural. Politekh. Inst. im. Kirova, Sverdlovek, USSR
Khimiya Geteroteiklichaskikh Boedinenii (1972), (4),
553-7
CODEN: KOSSAQ; ISSN: 0132-6244
JOURNAL
AUGUMGE: R. Kl.; Kusian
QI For diagram(s), see printed CA Issue.

For diagram(s), see printed CA Issue.

Five 2-phenazinecarboxamide 10-oxides (I, R = H, Et, R1 = H, Et, Ph,

p-MeOC6H4, RRI - (CH2CH2)20, n - 0) and 4 2-phenazinecsrboxamide 5,10-dioxides I (n = 1) were obtained in 46-86% yield. Polarog, of 36 I and previously obtained phenazinecarboxamides was reported. 30806-87-2 30806-88-3 30905-87-0 30905-73-8 30905-74-9 37648-78-5 37648-80-9 37648-82-3 RL: RCT (Resctant); RACT (Reactant or reagent) (polarog, of) 30806-87-2 CAPLUS Glycine, N-(2-phenazinylcsrbonyl)-, ethyl ester (SCI, 9CI) (CA INDEX NAME)

30806-88-3 CAPLUS Glycine, N-(2-phenszinylcsrbonyl)-, ethyl ester, dioxide (9CI) (CA INDEX NAME)

30905-67-0 CAPLUS L-Valine, N-(2-phenazinylcarbonyl)-, ethyl ester (9CI) (CA INDEX NAME) Absolute stereochemistry.

10905-73-6 CAPLUS L-Valine, N-(2-phenazinylcarbonyl)-, ethyl ester, dioxide (9CI) (CA INDEX NAME)

30806-89-4 RL: RCT (Rescient); RACT (Reactant or resgent) (pollrog. of) 30806-89-4 (APPLUS Glycine, N-(2-phenazinylcsrbonyl)-, dioxide (9CI) (CA INDEX NAME)

37648-67-2P 37648-71-8P 37648-75-2P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and polarog. of) 37648-67-2 CAPLUS 2-Phenazinecarboxamide, N.N.diethyl

-Phenazinecarboxamide, N,N-diethyl-, 10-oxide (9CI) (CA INDEX NAME)

37648-71-8 CAPLUS
2-Phenszinecsrboxamide, N,N-diethyl-, 5,10-dioxide (9CI) (CA INDEX NAME)

RN 37648-75-2 CAPLUS CN 2-Phenazinecarboxylic scid, ethyl ester, 10-oxide (9CI) (CA INDEX NAME)

RN 30905-74-9 CAPLUS CN L-Valine, N-(2-phenazinylcarbonyl)-, dioxide (9CI) (CA INDEX NAME)

Absolute stereochemistry.

37648-78-5 CAPLUS 2-Phenszinccsrboxylic acid, ethyl ester, 5,10-dioxide (9CI) (CA INDEX NAME)

37648-80-9 CAPLUS 2-Phenezinecarboxylic acid, ethyl ester (9CI) (CA INDEX NAME)

RN 37648-82-1 CAPLUS CN 2-Phenazinecarboxamide, N,N-diethyl- (9CI) (CA INDEX NAME)

Patent

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|-----|---------------------|--------|--------------|--------------------------|-----------|
| | | | | | |
| | DE 2144297 | A | 19720309 | DE 1971-2144297 | 19710903 |
| | DE 2144297 | C2 | 19621202 | | |
| | CH 553428 | A | 19740830 | CH 1970-13253 | 19700904 |
| | FR 2106207 | A5 | 19720428 | FR 1971-31379 | 19710830 |
| | US 3796576 | A | 19740312 | US 1971-176749 | 19710831 |
| | GB 1360046 | A | 19740717 | GB 1971-41053 | 19710902 |
| | BE 772142 | A1 | 19720303 | BE 1971-107759 | 19710903 |
| | JP 54003620 | B4 | 19790224 | JP 1971-68809 | 19710904 |
| | US 3875158 | A | 19750401 | US 1973-344815 | 19730326 |
| PRI | ORITY APPLN. INFO.: | | | CH 1970-13253 A | 19700904 |
| | | | | US 1971-176749 A | 19710831 |
| ΔD | Quinovalines, havin | a a fa | vorable redo | x potential and sdequate | solubilit |

Quinoxalines, having s favorable redox potential and adequate solubility in the dye blesch bath, carry in their 2 and 1-positions a -CU2 group linked to Br. C. O. S or N. They are synthesized from 1,2-dintrobenzene or o-nitroantline compds. by hydrogenstion to the 1,2-dintrobenzene or o-nitroantline compds. by hydrogenstion to the 1,2-dintrobenzene or followed by condensation with 1,2-diesrbonyl compds. in a N stmospheric The halogen in the 2,3-bis(bromomethyl) quinoxalines can resdily be exchanged by reaction with Levis bases. Thus, 2,3-bis(bromomethyl)-6,7-di(methoxy) quinoxaline is obtained by reduction of 1,2-dimethoxy-4,5-dinitrobenzene and condensation with BrRIGCOCCHSBr. Resection with [1, XICO] in aqueous EtON replaces the Br by ON, (2) KOAc by OAc, and (3) NsOMe by OMe. The quinoxalines are added to the blesch bath (1-100 mg/l.) or incorporated in a layer of the photog, material. 37966-43-1
RL: USES (Uses)
(photog, silver-dye bleach bath containing, for color processing)

IT

RL: USES (USES) (USES) (photog. silver-dye bleach bath containing, for color processing) 17966-43-1 CAPUS 6-Quinoxalinecsrboxylic scid, 2,3-bis[(scetyloxy)methyl]-, ethyl ester (9CI) (CA INDEX NAME)

L13 ANSWER 145 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
177:1434584 CAPLUS
77:34584
Anticancerous 2-substituted phenazine 5,10-dioxides
800/RCE:
800/RCE des usines chimiques de Rhone-Poulenc
Fr. CAM, 2 pp. Addn. to Pr. M 4745 (CA 69;67421f).
CODEN: PRIXISK
PATENT

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

PR 193 19650410 FR 1965-60633 19650618
FOR diagram(s), see printed CA Issue.

N.N'-carbonylddimidazole in DMF was treated with 2-carboxyphenazine
5,10-dioxide to give 2-imidazolocarbonylphenazine 5,10-dioxide.
1-Benzylpherazine was added, and the solution concentrated to give
2-(4-benzyl-1-piperazinyl). Similarly prepared were I [R =
([3-(dimethylamino)ethyl)amino], [(3-(dimethylamino)ropyl)amino],
(3-(4-pyrrolidinyl)ethyl)amino], [(3-(4-methyl-1-piperazinyl)]
13458-23-82 13458-27-07 13458-29-27
13458-23-82 13458-27-07 13458-29-27
Rh. (preparation) (preparation); PREP (Preparation)
(preparation of)
13458-23-82 CAPLUS
2-Phenazinecarboxamide, N-[2-(dimethylamino)ethyl]-, 5,10-dioxide (SCI, 9CI) (CA INDEX NAME)

13458-27-0 CAPLUS
2-Phenazinecarboxamide, N-{1-(dimethylamino)-2-methylpropyl}-,
5,10-dioxide (&CI, 9CI) (CA INDEX NAME)

its derivatives
Gordienko, L. 1, Rozum, Yu. S.; Prokopenko, V. P.
Klev. Tekhnol. Inst. Pishch. Prom., Klev. USSR
Elektrokhimiya (1971), 7(13), 1830-3
COENN: ELKOAK; 18SN: 042-8570
JOURNAL
Russian

DOCUMENT TYPE: LANGUAGE: AB Polarog.

PAGE 1-A ме~ (СН₂) 4 - О- С

PAGE 1-B

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

KIND DATE ---- 1971061 APPLICATION NO. US 1969-849600 US 1969-849600 PATENT NO. DATE

PATENT NO. KIND DATE APPLICATION NO. DATE

123 3585181 A 19710615 US 1969-849600 19690807

FOR diagram(s), see printed CA Issue.

B Hydroxy azonaphthalene Cu complex fiber reactive dyes (I, A = sulfo-1,2-naphthylenes, R = 2,3-dichloro-6-quinoxalinylcarbonyl), 4,6-dichloro-s-triexin-2-yl, R1 = H, SOSH, R2 = H, SOSH, R3 = H, SOSH, N3 = H, SOSH), useful for dyeing natural and regenerated cellulose, were prepared Thus, 2-(T-2(-[2,1-dichloro-6-quinoxalinylcarbonyl)methylaminolacetamidGomethyl)-2,8-dihydroxy-3,6-disulfo-1-naphthylamol-4-sulfo-1-naphthol 1:1 copper complex (I, A = sulfo-1-naphthylamol, R2 = H) vas prepared by condensing I

13458-29-2 CAPLUS 2-Phenazinecarboxamide, N-[2-(1-pyrrolidinyl)ethyl]-, 5,10-dioxide (8CI, 9CI) (CA INDEX NAME)

13458-30-5 CAPLUS 2-PhenazincarDoxamide, N-[3-(dimethylamino)propyl]-, 5,10-dioxide (aCI, 9CI) (CA INDEX NAME)

14559-63-8 CAPLUS
2-Phenazinecarboxamide, N-{2-(4-methyl-1-piperazinyl)ethyl}-, 5,10-dioxide
(BCI, 901) (CA INDEX NAME)

L13 ANSWER 146 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1972:419006 CAPLUS
DOCUMENT NUMBER: 77:19006
TITLE: 70:19006
Polarography and ir spectra of 2,2'-biphenazinyl and

(A = 4-sulf/-1, 2-naphthylene, R = H, R1 = R3 = SO3H, R2 = H) with 2,3-dichloroquinoxaline-6-carbonyl chloride at pH 7-8, and used to dye cotton fast blue shadas.
16207-38-89 16207-39-99 16265-96-69
16265-97-79 3311:1-5-89
RL: IMF (Industrial manufacture); PREP (Preparation) (preparation of)
16207-38-8 CAPLUS
Copper, [crihydrogen 3-[2-(2,3-dichloro-N-methyl-6-quinoxalinecarboxamido)acetamido)methyl]-4,6-dihydroxy-5-[(1-hydroxy-4-sulfo-2-naphthyl)azo]-2,7-naphthalenedisulfonato(2-)]- (SC1) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

16207-39-9 CAPLUS
Copper. [trihydrogen 3-[[7-[[2-(2,3-dichloro-N-methyl-6-quinoxalinearboxamido]acetamido]bsthyl]-2,8-dihydroxy-4-sulfo-1-naphthyl]azo]-4-hydroxy-1,5-naphthalenedisulfonato(2-)]- (BCI) (CA INDEX NAME)

PAGE 2-A

16265-96-6 CAPLUS

Copper, [trihydrogen]-[[7-[[2-(2,3-dichloro-N-methyl-6-quinoxalinecarboxamido]acetamido]bethyl]-2,8-dihydroxy-6-sulfo-1-naphthyl]azo]-4-hydroxy-1,5-naphthalenedisulfonato(2-)]- (SCI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

●4 H+

33111-15-8 CAPLUS
COpper, [dihydrogen 3-[[[[2-(2,3-dichloro-N-methyl-6quinoxalinearboxamido]acetamido]methyl]-2-hydroxy-3-methanesulfonamido-1naphthyl]szo]-4-hydroxy-1,5-naphthalenedisulfonato(2-)]- (SCI) (CA INDEX
RAMES)

PAGE 2-A

16265-97-7 CAPLUS
COpper, [[tetrahydrogen 6-[[2-(2,3-dichloro-N-methyl-6quinoxalinecarboxamido)acetamido]methyl]-3,3',5-trihydroxy-4,4'-azodi-2,7naphthalenedisulfonato](2-)]- (8CI) (CA INDEX NAME)

PAGE 1-A -035

$$\begin{array}{c|c} & & & & & & \\ & & & & & \\ D1-CH_2-NH-C-CH_2-N-C & & & & \\ & & & & & \\ \end{array}$$

●2 H+

LI3 ANSWER 148 OF 181
ACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE:
AUTHOR(5):
CORPORATE SOURCE:
SOURCE:
AUTHOR(5):
CORPORATE SOURCE:
SOURCE:
CORPORATE SOURCE:
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
AUTHOR(5):
CORPORATE SOURCE:
SOUR

L13 ANSWER 149 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE:
TITLE:
TAYLOROGY:
TAYLOROGY:
TOTALE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

DATE PATENT NO. A C3 B2 DATE APPLICATION NO. DE 2010280 DE 2010280 DE 2010280 CH 508226 US 3656953 FR 2034876 BE 747252 NL 7003551 NL 167523 GE 1299402 SU 383336 JP 49010054 19700924 DE 1970-2010280 19700305 19791115 19791115 19790322 19710531 19720418 19701218 19700914 19700915 19810716 19811216 19721213 CH 1969-508226 US 1970-16207 FR 1970-8444 BE 1970-747252 NL 1970-3551 19690313 19700303 19700310 19700312

| NL 167523 C 19811216 | GB 1279402 A 19721213 GB 1370-1299402 19700312 | GB 1279402 A 19721213 GB 1370-1299402 19700312 | GB 1279-1416280 19700312 | GB 1279-1416280 19700313 | GB 1279-1416280 19700313 | GB 1279-1416280 | GB 12700313 | GB 1279-14162 A 19700313 | GB 127900313 | GB 12790031 | GB 127

32388-06-0 CAPLUS

30806-87-2 CAPLUS Glycine, N-(2-phenazinylcarbonyl)-, ethyl ester (&CI, 9CI) (CA INDEX

30806-88-3 CAPLUS Glycine, N-(2-phenazinylcarbonyl)-, ethyl ester, dioxide (9CI) (CA INDEX NAME)

30806-89-4 CAPLUS Glycine, N-(2-phenazinylcarbonyl)-, dioxide (9CI) (CA INDEX NAME)

- NH- CH2- CO2H

30905-67-0 CAPLUS L-Valine, N-(2-phenazinylcarbonyl)-, ethyl ester (9CI) (CA INDEX NAME) Absolute stereochemistry.

30905-73-8 CAPLUS L-Valine, N-(2-phenazinylcarbonyl)-, ethyl ester, dioxide (9CI) (CA INDEX NAME)

Absolute stereochemistry.

6-Quinoxalinecarboxylic acid, 2,3-dimethyl-, ethyl ester (6CI, 8CI, 9CI) (CA INDEX NAME)

32388-08-2 CAPLUS 6-Quinoxalinecarboxamide, N,N-diethyl-2,3-dimethyl- (8CI, 9CI) (CA INDEX NAME)

32388-09-3 CAPLUS 6-Quinoxalinecarboxamide, N.N-diethyl-2,3-diphenyl- (&CI, 9CI) (CA INDEX NAME)

L13 ANSWER 150 OF 101 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1971:87918 CAPLUS
DOCUMENT NUMBER: 74:87918
TITLE: N-Oxides of N-phenazinoyl derivatives of some

Q-amino acids Batulina, R. Kh.; Pushkareva, Z. V.; Konyukhov, V. N.; Bobarykina, K. Yu.; Platonova, G. N. Ural. Politckh. Inst. im. Kirova, Sverdlovsk, USSR Khimiko-Parmateevticheskii Zhurnal (1970). 4(11). AUTHOR (S) :

CORPORATE SOURCE: SOURCE: 18-22

CODEN: KHFZAN; ISSN: 0023-1134 Journal

DOCUMENT TYPE:

DOCUMENT TYPS: Journal
LANGUAGE:
Russian
GI For diagram(s), see printed CA Issue.
AB 1-Phenazinecarboxylic acid was obtained from condensation of
o-H2NC6H4CO2H and PhNO2. 2-Phenazinecarboxylic acid (I) was
prepared by oxidative condensation of p-toluidine with PhNO2. I as the acid
chloride reacted with amino acid Et esters to give II. III and IV were
obtained from II (R = iso-Pr) by oxidation with 30 H2O2 in HOAc.

IT 10806-87-29 10806-88-39 10806-89-49
30905-67-09 10905-73-89 30905-74-99
RL: SPN (Symthetic preparation); PREP (Preparation)
(preparation of)

30905-74-9 CAPLUS L-Valine, N-(2-phenazinylcarbonyl)-, dioxide (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L13 ANSWER 151 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1970:521552 CAPLUS
DOCUMENT NUMBER: 73:121552
PATENT ASSIGNEE(S): Farbenfabriken Bayer A.-O.
SOURCE: Fr. Demande, 12 pp.
CODEN: FRXXBL
DOCUMENT TYPE: Patent
LANGUAGE: Proch

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. APPLICATION NO. KIND DATE FR 1969-19221 GB DE FR 2011515 GB 1262583 19700306 19690610

FR 201515 19 19700306 PR 1989-1921 19890610 GB 1262563 GB 1262563 DE 08 1980610 GB 201517 APPLM. INFO.:

OB 201517 APPLM. INFO.:

OB 201518 DE 01 1980610 DE 1980610

PAGE 2-A

2 D1-803-

●2 Na+

RN 29116-77-6 CAPLUS
CN Copper, [trihydrogen [[2-{2,3-dichloro-N-methyl-6-quinoxalinecarboxamido]ethyl]methylsulfamoyl]phthalocyaninetrisulfonato(2-)]-, trisodium salt (SCI) (CA INDEX NAME)

OTHER SOURCE(S):

MARPAT 72:100749
GI For diagram(s), see printed CA Issue.

AB The title compds. (I or II) were prepared by treating an amine with 6-(chlorocarbony)-2,3-dichloroquinoline (III) or III derivs. The compds. inhibit acid secretion. Thus, 0.048 mole CHANNACH2OH in 2.3 ml, dioxane added to 0.023 mole III in 52 ml dioxane gave 6-(2-hydroxyethyl)carbamoyl)-2,3-dichloroquinoxaline, m. 181-3* (McCN). Similarly were prepared 6-(diethylcarbamoyl)-2,3-dichloroquinoxaline, m. 73-5*, 6-((2-methoxyethyl)-carbamoyl)-2,3-dichloroquinoxaline, m. 141-2*, and the following I (X = H) (R, R1, and m.p. given): CH2CH2OAC, H. 161.5-3.5*; Me, H. 213-15*; Me, Me, Me, 138-9*; Et. H. 191-6*; Et. Et. 73-5*; Pr. H. 158-60*; CH2CH2OH, Me, 128-30*; CH3CH2OH, H. 157-9*; CH2CH2OH, Me, 128-30*; CH3CH2OH, H. 157-9*; CH2CH2OH, M. 180-2*; CH2CH2OH, M. 181-2*; CH2CH2OH, M. 181-2*; CH2CH2OH, M. 181-2*; CH2CH2OH, H. 185-6*; CH2CH3OH, M. 180-2*; CH2CH2OH, M. 181-2*; CH2CH2OH, H. 181-6*; CH2CH2OH, M. 180-2*; CH2CH2OH, M. 181-2*; CH2CH2OH, M. 180-6*; CH2CH2OH, M. 180-6*;

PAGE 2-A

3 D1-503-

●3 Na+

L13 ANSWER 152 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1970:100749 CAPLUS
OCCUMENT NUMBER: 72:100749
TITLE: Quinoxalines useful in treatment of peptic ulcers
INVENTOR(S): Bolhofer, William A.; Beldwin, John J.
Merck and Co., Inc.
SOURCE: S. African, 25 pp.
CODEN: SFYAMB
DOCUMENT TYPE: Patent
English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|----|----------------------|------|----------|-----------------|----------|
| | | | | | |
| | ZA 6707613 | | 19690619 | | |
| | DB 1695532 | | | DE | |
| | PR 1588778 | | | FR | |
| | FR 7331 | | | FR | |
| | GB 1180249 | | | GB | |
| | US 3510487 | | 19700505 | US | 19661228 |
| | US 3655894 | | 19720411 | us | 19690828 |
| PR | IORITY APPLN. INFO.: | | | US | 19661228 |
| | | | | | |

RN 26773-14-8 CAPLUS CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-methoxyethyl)- (8CI) (CA INDEX NAME)

RN 26773-17-1 CAPLUS CN 6-Quinoxalinecarboxamide, 2,3,7-trichloro-N-(2-hydroxyethyl)- (8CI) (CA INDEX NAMS)

RN 26773-18-2 CAPLUS
CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-[2-(dimethylamino)ethyl]- (8CI)
(CA INDEX NAME)

RN 26773-19-3 CAPLUS CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-morpholinoethyl)- (8CI) (CA INDEX NAME)

$$\bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-NH-C} \bigcap_{0}^{N-CH_{2}-NH-C}$$

RN 26773-21-7 CAPLUS CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-[2-(dimethylamino)ethyl]-, hydrochloride (ECI) (CA INDEX NAME)

Ox HCl

RN 26773-22-8 CAPLUS CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-hydroxyethyl)-, acetate (ester) (8CT) (CA INDEX NAME)

RN 26773-25-1 CAPLUS CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-ethyl- (8CI, 9CI) (CA INDEX NAME)

RN 26773-26-2 CAPLUS CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N,N-bis(2-hydroxyethyl)- (8CI) (CA INDEX NAME)

RN 26773-27-3 CAPLUS
CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-hydroxyethyl)-N-methyl- (8CI)
(CA INDEX NAME)

RN 26773-28-4 CAPLUS
CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-hydroxy-1-methylethyl)- (@CI)
(CA INDEX NAME)

RN 26773-29-5 CAPLUS CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-hydroxypropyl)- (8CI) (CA INDEX NAME)

RN 26773-30-8 CAPLUS
CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-hydroxy-2-methylpropyl)- (8CI)
(CA INDEX NAME)

RN 26773-31-9 CAPLUS
CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(3-hydroxypropyl)- (8CI) (CA INDEX NAME)

RN 26773-32-0 CAPLUS CN 6-Quinoxilinecarboxamide, 2,3-dichloro-N-(2-chloroethyl)- (8CI, 9CI) (CA INDEX NAME)

RN 26840-63-1 CAPLUS CN 6-Quinoxalinecarboxemide, 2,3-dichloro-N-(tetrahydrofurfuryl)- (8CI) (CA INDEX NAME)

RN 26840-68-6 CAPLUS
CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2,3-dihydroxypropyl)- (8CI) (CA INDEX NAME)

RN 26840-73-3 CAPLUS
CN 6-Quinoxalinecarboxamide, N-(2-acetamidoethyl)-2,3-dichloro- (8CI) (CA INDEX NAME)

RN 26840-74-4 CAPLUS
CN 6-Quinoxalinecerboxamide, 2,3-dichloro-N-(β-hydroxyphenethyl)- (SCI)
(CA INDEX NAME)

RN 26840-75-5 CAPLUS
CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-[4-(diethylamino)butyl]-,
monohydrochloride (8CI) (CA INDEX NAME)

• HC1

RN 26840-76-6 CAPLUS
CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-[2-(diethylemino)ethyl]monohydrochloride (8CI) (CA INDEX NAME)

$$\underset{\text{Et}_{2}\text{N}-\text{CH}_{2}-\text{CH}_{2}-\text{NH}-\text{C}}{\overset{\overset{\circ}{\text{N}}}{\overset{\circ}{\text{N}}-\text{C1}}} \overset{\circ}{\underset{\text{C1}}{\text{N}}}$$

• HC1

RN 26840-77-7 CAPLUS
CN 6-Quinoxalinecarboxamide, 2,3-dichloro-N-[3-(dimethylamino)propyl]-,
monohydrochloride (sCI) (CA INDEX NAME)

• HC1

RN 26840-78-8 CAPLUS

6-Quinoxalinecarboxamide, 2,3-dichloro-N-(2-morpholinoethyl)-, monohydrochloride (8CI) (CA INDEX NAME)

● HC1

26887-34-3 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N,N-diethyl- (8CI, 9CI) (CA INDEX NAME)

26887-35-4 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-propyl- (8CI) (CA INDEX NAME)

26921-20-0 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dichloro-N-2-propenyl- (9CI) (CA INDEX NAME)

LI3 ANSWER 153 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1970:70530 CAPLUS
DOCUMENT NUMBER: 72:70530
TITLE: Preparation and laboratory evaluation of cellulose-based ion permedicative membranes
AUTHOR(S): Sueszer, A.; Bandel, E.; Flitman, M.
CORPORATE SOURCE: Negev Inst. Arid Zone Res., Beer-Sheva, Israel

stirring followed by 10 parts NaCl, the precipitated dye filtered, washed with

solution of 24 parts NaCl, 0.37 part NaHSO4, and 6 parts VI in 240 parts H2O, and dried at room temperature Similarly, other dyes were prepared (reactants given): II, I (Y = CHZCHMe, Z = H) (VII), III; II, VII. 5-cyano-2,4.6-trichloropyrimidine: II, VII, 2,3-dichloro-6-quinoxalinecarbonyl chloride; I (Y = CHZCHZ, Z = CHZCHZOH), III, mixed with IV. 25138-46-49 25238-47-59
RL: INF (Industrial manufacture); PREP (Preparation) (preparation of

IT

(preparation of)

2523a-6-4 CAPUS

2523a-6-4-6 CAPUS

2523a-6-4-6 CAPUS

quinoxalinecarboxamido) ethyl] sulfamoyl] anilino) -9,10-dihydro-9,10-dioxo-(8C1) (CA INDEX NAME)

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25238-47-5 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[m-[[2-(2,3-dichloro-6-quinoxalinecarboxamido]propyl]sulfamoyl]anilino]-9,10-dihydro-9,10-dioxo-(8CI) (CA INDEX NAME)

SOURCE: Desalination (1969), 7(1), 47-50
CODEN: DSLNAN; ISSN: 0011-9164
DOCUMENT TYPE: Journal
LANGUAGE: English
AS Ion permselective membranes were prepared by treating cellophane membranes
with trichloropyrisidine reactive dyes, i.e. Drimarene Black Z-BL or
Reaction Yellow RL, or dichloroquinoxaline reactive dyes, i.e. N-(3-d
imethylaminopropyl)-3,3-dichloro-6-quinoxalinecar-boxamide. Promising
results were obtained when electrodialytic desalination stacks prepared with
the membranes were used in the desalination of water.

IT

the membranes were used in the desalination of water.
24604-56-6
RL: OCCU (Occurrence)
(reaction products, with cellophane)
24604-56-6
CAPILUS
6-Quinoxalinecarboxamide, 2,3-dichloro-N-[3-(dimethylamino)propyl]- (SCI)
(CA INDEX NAME)

L13 ANSWER 154 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
11:11452
Reactive dyes
FITTLE:
PATENT ASSIGNEE(S):
FOURCE:
CODEN: FRXXAX
PATENT
PATENT
CODEN: FRXXAX
PATENT
PATENT DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DATE APPLICATION NO. DATE FR 1531271 DB 1619594 GB 1130395 GB 1180395 19670718 FR 1967-114662 DE 19680628 GB US 3489502 PRIORITY APPLN. INFO.: 19700113

For diagram(s), see printed CA Issue.

Compds, of the general formula I, where Y is an alkylene or arylene radiced. The see that the set of the general formula I, where Y is an alkylene or arylene radiced. Thus, a see that the set of the general formula I, where Y is an alkylene or arylene radiced. Thus, a see that the set of th

PAGE 1-A

PAGE 2-A

L13 ANSWER 155 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1595:492648 CAPLUS
TITLE: 71:92648 TAPLUS
INVENTOR(5): Anthraquinone fiber-reactive dyes
Harms, Wolfgang: Gehrke, Gunter; Hohmann, Walter;
Bien, Hens S.
PATENT ASSIGNEE(8): Farbenfabriken Bayer A.-O.
BOCUMENT TYPE: Patent

DOCUMENT TYPE: Patent

Patent English

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|-----------------------|---------|--------------|-----------------|----------|
| PAIBNI NO. | | DATE | APPLICATION NO. | |
| | | | | |
| GB 1147297 | | 19690402 | GB 1967-54744 | 19671201 |
| DE 1644612 | | | DE | |
| FR 1551267 | | | FR | |
| RIORITY APPLN. INFO.: | | | DE | 19661207 |
| For diagram(a) . se | e print | ed CA Leaun. | | |

Compde. of the general formula I where X is a group containing at least one reactive halogen, are water soluble yellow-green to green reactive dyes for

PhCl is removed by seration, the mixture clarified and salted to precipitate

H, X = CCO) a yellowish green dye. Similarly other yellowish green I (Y =
OH) are prepared (R and X given): H, CSO2, 2-methoxyeutlonyl-4-methyl-5chloro-6-pyrimidinyl (2): H, 4-ZNHCSH4CO; H, 3-ZNHCSH4SO2; H,
2,4-difluoro-5-chloro-6-pyrimidinyl; H, CLC12CO; H, MeMCHACO; H,
CSO2NMCCH2CO; H, CCONMCCH2CO; H, BCCH2CHECO; H, 1,4-dichloro-6phthalarimecarbonyl; H, 4,6-dichloro-s-triarin-2-yl(O'); H,
2,4-5-trichloro-6-pyrimidinyl; H, 4-chloro-6-methoxy-a-triazin-2-yl,
Similarly other green I were prepared (R, X, and Y given): H, CCO, NH2; H,
CSO2, NH2; H, CCO, Cl; H, CSO2, Cl; H, O', Cl; H, CCO, Br; H, CCO,
NH2; H, CCO, and blue-green II (X = SO2),
23945-59-4 23945-02-13P

RLi IMF (Industrial manufacture); PREP (Preparation)
(preparation of)
23945-58-4 CAPLUS

T-Cluenesulfonic acid, 6,6'-[[5-{2-(2,3-dichloro-N-methyl-6quinoxalinocarboxamido) acetamido]-8-hydroxy-1,4anthraquinonylene]dimino]di-, disodium salt (aCI) (CA INDEX NAME)

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L13 ANSWER 156 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NAMBER: 1998-442641 CAPLUS
TITLE: Anthony of the receive dyes
INVENTOR(S): Leister, Heinrich
SOURCE: COPYRIGHT 2006 ACS on STN
112641 CAPLUS
T1:2641 CA

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|----|--------------------|------|----------|-----------------|----------|
| | | | | | |
| | GB 1147110 | | 19690402 | GB 1967-53799 | 19671127 |
| | DR 1644611 | | | DE | |
| | FR 1546177 | | | FR | |
| | US 3828040 | | 19740806 | US 1970-64007 | 19700717 |
| 10 | RITY APPLN. INFO.: | | | DE | 19661130 |
| | | | | | |

PR 1546177

PR US 3822000

1930800

19740806

19 1970-64007

1970717

19651130

POT diagram(a), see printed CA Issue.

I, where X is a group containing a reactive halogen, and II, where R is sulfophenyl or sulfato, are water-soluble yellowish green to bluish green dyes for cotton. I are prepared by reacting the appropriate

S-amino-1,4-bisanilinoenthraquinone with a reactant containing ≥2

reactive halogen atoms. A mixture of 15 parts 5-aminoquinizarin, 100 parts

P-McCGRAMIN, 8 parts H3B03, and 6.3 parts concentrated HC1 is treated at 75° with 2 parts ≥7 with 100 parts has, treated at 55° with 15 parts powdered KOR, the melt aersted, cooled to 65°, and treated with 100 parts MoOH to give I (X = R = B) (III), green needles. III (20 parts) is added during 3 hrs. to a mixture of 48 parts 200 oleum and 52 parts concentrated H2S04. The mixture is treated with 30 aparts 200 oleum, stirred until disulfonation is complete, and treated with 300 parts ice to precipitate I (X = R, R = S018) (IV). A ution of 8

parts 18 salt of IV in 150 parts H20 is heated at 80.5° with 7

parts 24.5,6-tetrachloropyrimidine (V) for 12 hrs. while the pH is kept at 5.5-7.0 by dropwise addition of N aqueous NaOH, excess V boiled off, and the green dye I (X = 2,4-5-trichloro-6-pyrimidinyl, R = S018n) (VX) is salted out. The following I (R = S03Na) are similarly prepared from IV (X and shade on cotton given): 4-chloro-6-tenthoxy-a-triaxin-2-yl, green; 2,3-dichloro-6-quinoxa-linylcarbonyl, green; 2,3-dichloro-8-quinoxalinylsulfonyl, bluish green; 4,6-dichloro-6-quinoxa-1-yprepared (X given): p-03M6H4CO (VI); m-03M6H6CO (VI); m-03M6H6CO (VI); m-03M6H6CO (VI); m-03M6H6CO (VI); m-03M6H6CO (VII). A solution of VI (from 55 parts Na salt of IV) in 855 parts Ros and solution of VI (from 55 parts Na salt of IV) in 855 parts Ros and solution of VI (from 55 parts Na salt of IV); also parts Na salt of IV) in 850 parts Ros and solution of VI (from 55 parts Na salt of IV); also parts Ros and solution of VI (from 55 parts Na salt of IV); also parts Ros and

PAGE 2-A

23946-02-3 CAPLUS
m-Toluenesulforic acid, 6,6'-{[5-[2-(2,3-dichloro-N-methyl-6quinoxalinesulfonamido]acetamido]-8-hydroxy-1,4anthraquinonylene]dimino]di-, dieodium salt (8CI) (CA INDEX NAME)

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PAGE 1-A

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L13 ANSWER 157 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOJUMENT NUMBER:
TITLE:
TITLE:
Piber reactive azo dyes
INVENTOR(S):
PATENT ASSIGNEE(S):
Farenfabriken Bayer A.-G.

PATENT ASSIGNER (8) : SOURCE: Brit., 10 pp. CODEN: BRXXAA

Patent English DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. GB 1100228 19681009 GB 1967-40454 19670905
DE 1644170 DE FR
FS 1538115 FR
PRIORITY APPLM. INFO: DE 19660910
GI FOR diagram(s), see printed CA Issue.
AB The moist paste of the monoazo compound obtained by diazotism 31.3 parts 4,3-H3N(MO38)C68102H3NHs and coupling with 46.1 parts 1,5,4,7(MO38)2C10814(NM2)2 is slurried in 500 part H20, treated with 200 parts 204
NM40H and 80 parts CuS04.5H20, the mixture heated at 90-5° until the red color has disappeared, treated with Na2S to precipitate Cu2S and the

red color has disappeared, treated with Na28 to precipitate Cu28 and the solution of the resultant triaxole (I, ONH2) diaxotized and coupled (acid) with 34.7 parts 1,3,7-No(Mos)SC10MSNN2. The moist paste of the azotriazole derivative is stirred into 1000 parts H20, treated with 12 parts NaNO2, poured into a solution of 12.5 parts concentrated M2804 in 200 parts ice-water, and stirred overnight, and salted to give 6,2,8,1-NO38-(NO)2C10M4N: NO. The moist cake is dissolved in 500 parts H20 at 40°, treated with 40 parts 2,3-dichloroquinoxaline-6-carboxylic acid chloride, maintained at pH 5.7 for 10 hrs. by addition of Na2OO3, added to 1000 parts H20. clarified with charcoal, and salted. The moist cake is suspended in 4000 parts ice-water and added to a solution of 40 parts Cu504-5802 in 400 parts H20, kept at pH 7 by addition of aqueous NaOH, and the dark red solution mixed dropwise with 250 parts

34 H202 (turns blue) and salted to give II, a clear blue, fiber reactive dye. 22873-61-6P RE: 1MF (industrial manufacture); PREP (Preparation)

RL: IMF (Industrial manufacture); PREP (Preparation) (preparation of) 22873-61-6 CAPLUS Copper, (Estrahydrogen 2-[α-(2,3-dichloro-N-methyl-6-quinoxalinecarboxamido)-6-eulfo-m-tolyl]-7-[(2,8-dihydroxy-6-aulfo-1-naphthyl)axo]-6-hydroxy-2H-naphtho[1,2-d]triazole-5,9-disulfonato(2-)]-(SCI) (CA INDEX INDEX)

19625-39-9P
RL: PORM (Formation, nonpreparative); PREP (Preparation)
(Commation of)
19625-39-9 CAPJUS
Copper, ju-[dihydrogen 5,8-dihydroxy-6,7-quinoxalinedicarboxylato(2-)]]bis(dihydrogen 5,8-dihydroxy-6,7-quinoxalinedicarboxylato)di-,hexaethyl ester (8C1) (CA INDEX NAWS)

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ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

ANSWER 158 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN

SSSION NUMBER: 1968:413168 CAPLUS
69:13368 CAPLUS
69:13368 CAPLUS
69:1368 CAPLUS
60:1368 AUTHOR(S): CORPORATE SOURCE: SOURCE:

CODEN: BCSJA8; ISSN: 0009-2673

DOCUMENT TYPE: Journal
LANGUAGE:
AB The solubility, ionization constant, and chelating behavior of
dihydroxyquinoxaline, mainly 5,8-dihydroxyquinoxaline (1) derivs., were
measured. The introduction of a hydroxyl group into quinoxaline greatly
lowers the solubility The order of the decreasing solubility of
2,3-disubstituted 1
in H20 (at 20°) is: O(CH2)2-OBt < H < IH < Me < Ph < Eto < Cl < SH
< Etc. For I the introduction of electron-repelling groups into
2,3-positions raises both the pKNH and pKOH values, while the instruction
of electron-attracting groups into the same positions lowers both pKNH and
pKOH. I and its derive. form colored precipitate with metal ions, but some
derives, which have pKNH values lower than zero fail to show any precipitation
or

coloration. 6,7-Dihydroxyquinoxaline forms precipitate, and the precipitation

distinctively with Cr(III) or Fe(II). The composition and stability constant of Cu(II) chelates of I and its derivs. in a dioxane-H2O (10:90 by volume) solution were studied spectrometrically. The Cu chelate of I had a metal:-ligand ratio of 1:1 and a log K value of 6.28 at pH 4.0. 27 references.

2427-91-0
RL: PRP (Properties)
(chelation properties and ionization and solubility of)
2427-91-0 CAPLUS
6.7-Quinoxelinedicarboxylic acid, 5.8-dihydroxy-, diethyl ester (7CI, 8CI)
(CA INDEX NAME)

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2427-91-0DP, 6,7-Quinoxalinedicarboxylic acid, 5,8-dihydroxy-, diethyl ester, copper complex (RL: SPM (Synthetic preparation); PREP (Preparation) (preparation of) 2427-91-0 CAPLUS 6,7-Quinoxalinedicarboxylic acid, 5,8-dihydroxy-, diethyl ester (7CI, 8CI) (CA INDEX NAME)

L13 ANSWER 159 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 156:21412 CAPLUS
DOCUMENT NUMBER: 68:21412 CAPLUS
FOR COMMENT NUMBER: 156:21412 CAPLUS
OLINOXALINE and monosubstituted quinoxalines
AUTHOR(S): Brignell, Peter J.; Karritzky, Alan R.; Reavill, Roger
Brignell, Peter J.; Serfield, A. A.
DOCHOGATE SOURCE: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: Journal of the Chemical Society (Section) B: Physical Organic (1967), (11), 1241-3
COUNGENT TYPS: JOURNAL ORGANIC (1967), (11), 1

L13 ANSWER 160 OF 181 CAPLUS COPYRIGHT 2006 ACS On STN

1967:500993 CAPLUS 67:100993 Metallized ato dyes containing 2,3-dichloroquinoxaline-6-carbonylamino groups Jeeger, Horst; Gerlach, Klaus Farbenfabriken Bayer A.-G. Fr. 9 no. INVENTOR(S):

PATENT ASSIGNEE (S) : SOURCE : Fr., 9 pp. CODEN: FRXXAK Patent French

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

KIND DATE APPLICATION NO. PATENT NO. DATE

PATENT NO. KIND DATE APPLICATION NO. DATE

PRI 1470126 19470217 FR 1966-51204 19460225

PRIORITY APPLM. INFO.:

OI For diagram(s), see printed CA Issue.

A I. II. and III are blue dyes for cotton. Thus, 27.6 parts
2,3.5-NO(NOSS) (MeNNICHIZCONN)(C612NN2 is diszotized and coupled with 34.1
parts 8,3.6,1-182(NOSS)) 2610HOHOH, to give an azo dye which is dissolved in 1000 parts water at 45°, treated with a solution of 24.9 parts CUSO4 at pR 5-6 (Na2CO3), and treated at 45° with 36.1 part
2,3-dichloroquinoxaline-6-carboxylic acid chloride (OC1) at pH 5-6 (Na2CO3) to give II. dark powder, which gives a violat-blue aqueous solution and reddish blue shades on cotton. Also prepared are (color on cotton given):

II 18246-84-79 16520-32-49
RI: IMY (Industrial manufacture); PREP (Preparation)
(preparation of)
RN 18246-84-7 CABUS
COpper, [trihydrogen 4-mino-6-[{5-[2-(2,3-dichloro-N-methyl-6-quinoxal inecarboxamido) acetamido)-2-hydroxy-3-sulfophenyl]azo]-5-hydroxy-1,3-naphthalenedisulfomato(2-)]- (SCI) (CA INDEX NAME)

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L13 ANSWER 161 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1967:482944 CAPLUS
DOCUMENT NUMBER: 7:622944
TITLE: 7:62294
Fiber-reactive dyes
Jaeger, Horst; Schuendehuette, Karl H.; Machatzke, Heinz
PATENT ASSIGNEE(S): Ferbenfabriken Bayer A.-G.

PATENT ASSIGNEE(S): SOURCE:

Pr., 7 pp. CODEN: FRXXAK Patent Prench

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. FR 1966-56034 DE GB DD PATENT NO. KIND DATE DATE FR 1474432 19670324 FR 1966-56034 19660401
DE 1544516 DE 09 1056033 09 PRIORITY APPLAY. INFO: DD 19650403
DD 1043gram(s), see printed CA Issue.
AB Blue monozo dyes containing the reactive group 0 are prepared by treatment of

copperized aso dye with CICHZCONNCH2ON (I) followed by MeNH2 and finally with 2,3-dichloroquinoxaline-6-carboxylic acid chloride (II). For example, a solution of 63.2 parts Cu complex of 2,8,3,6,1 (Mo)2 (HOS)22C10H3N:NCH05(OH)8OH+2,1,4 in 450 parts ice-cold 96% H2SO4 is stirred in an ice bath, treated with 18.5 parts finely ground I, stirred for 12 hrs. at 10-15%, and poured into 1500 parts of ice. The product is precipitated by addition of 75 parts NaCl, filtered, redissolved in

parts water, adjusted to pH 7, and repptd. with NaCl. The wet paste is atirred with 100 parts 35% MeNN2 and 100 parts water for 24-48 hrs. at ambient temperature, adjusted to pH 5 with concentrated HCl, and filtered. A solution of

O.1 mole of this product in 400 parts water at 30-40° at pH 7 is attracted with a suspension of 28 parts II together with a solution of Na2CO3 to keep pH 7-8, precipitated with KCl, filtered, and dried to give 1000

111 (M = Y = SO3H, X = Z = H, R = OR), dyeing blue shades. Similarly, III (Z = SO3H) are prepared (R, M, X, and Y given): OH, H, SO3H, H; MeSO2NH, H, H, CH, H, R, SO3H. 1V is prepared similarly.

16207-38-87 16207-39-97 16285-98-689

16265-97-79 33111-15-89, Copper, [dihydrogen 3 - [[[2-2], 3-dichloro-l-mechtyl-6-yulnoxalinecarboxamido] acetamido] methyl]-

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CAPLUS le320-32-4 CAPLUS
Copper, [trihydrogen 4-mmino-6-[[6-[2-(2,3-dichloro-N-methyl-6-quinoxalinecarboxamido) acetamido]-2-hydroxy-4-sulfo-1-naphthyllazo]-5-hydroxy-1,3-naphthalenedisulfonato[2-]- (SCI) (CA INDEX NAME)

PAGE 1-A

2-hydroxy-8-methanesulfonamido-1-naphthyl)azo]-4-hydroxy-1,5naphthalenedisulfonato(2-)]RL: IMF (Industrial manufacture); PREP (Preparation)
(preparation of)
16207-18-8 CAPLUS
Copper, [trihydrogen 3-[2-(2,3-dichloro-N-methyl-6quinoxalinecarboxamido]acetamido]methyl]-4,6-dihydroxy-5-[(1-hydroxy-4sulfo-2-naphthyl)azo]-2,7-naphthalenedisulfonato(2-)]- (SCI) (CA INDEX
NAME)

PAGE 1-A

16207-39-9 CAPLUS
Copper, [trihydrogen 3-[[7-[[2-(2,3-dichloro-N-methyl-6quinoxalinearboxamido]acetamido]acthyl]-2,8-dihydroxy-4-sulfo-1naphthyllazo]-4-hydroxy-1,5-naphthalenedisulfonato(2-)]- (BCI) (CA INDEX
NAME)

PAGE 2-A

16265-96-6 CAPLUS
Copper, [trihydrogen 3-[[7-[[2-(2,3-dichloro-N-methyl-6quinoxalinearboxamido]acetamido]ecthyl]-2,8-dihydroxy-6-sulfo-1naphthyllazo]-4-hydroxy-1,5-naphthalenedisulfonato(2-)]- (SCI) (CA INDEX
RAME)

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PAGE 2-A

●4 H+

33111-15-8 CAPLUS
Copper, [dihydrogen 3-{[[[2-(2,3-dichloro-N-methyl-6quinoxalincarboxamido]acetamido]bethyl]-2-hydroxy-8-methanesulfonamido-1naphthyl]szo]-4-hydroxy-1,5-naphthalenedisulfonato(2-)]- (8CI) (CA INDEX
NAME)

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RN 16265-97-7 CAPLUS CD Copper, [[tetrahydrogen 6-[[2-(2,3-dichloro-N-methyl-6-quinoxalinecarboxamido]acetamido]methyl]-3,3',5-trihydroxy-4,4'-azodi-2,7-naphthalenedisulfonato](2-)]- (SCI) (CA INDEX NAME)

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●2 H+

L13 ANSWER 162 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE:
STABLIZED (First Stabilized fiber-reactive dyes
Kisse, Erik
du Pent de Nemours, E. I., and Co.
U.S., 9 pp.
CODEN: USKXAM
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC, NUM. COUNT:
FAMILY ACC, NUM. COUNT:
THORMATION:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATS APPLICATION NO. DATE

193131797 19470411 US 1963-25056 1953017

For diagram(s), see printed CA Issue.
2,3-Dichloroquinoxaline dyee for cotton were prepared and stabilized against hydrolysis by formulation with NaSSO4 or KSSO4. Thus, 30 parts
2,3-dichloro-6-quinoxalinecerbonyl chloride (QCI) was added to a solution of 42 parts 4,8,2-(MOSS)2C10HSMR2 → 3-McC6H4NR2 in 700 parts H2O at 40° and pN 7.5-8, the mixture stirred overnight and salted with 10 parts NaSSO4 to give a light yellow powder containing .appxx.824 I, 11% H2O, and 7% NaSSO4. Similarly, other amino dyes were acylated with QCI and the resultant amides salted (or blended) with NaSSO4 or KSSO4 (amino dye used and shade of product on cotton given): 4-(3-sulto-5-aminophenylaxo) derivative of 1-(2,5-dichloro-4-sulfophenyl)-3-methyl-5-pyrezolone (II), greenish yellow [also prepared by coupling 2,4-H2N(QNN)C6H3SO3N with II]; 1:1 Cu

complex of 2.5.7.6-H2N(HO)(H03S)ClOH4H:NC6H3(OH)SO3H-2.5, rubine; III, blue; cupe(SO3Na)2.5(SO2NH21).4SO2NNC6H3(NH2)SO3Na-3.4 (Pc = Phthalocyanine), turquoiae; 3.4-AcnN[2.4-R03S(a-H03SCGH4N:N)CGH3N:N]CGH3NH2, -. A similar dye was prepared by acylating 1,8.3.6,7-H2N(HO)(NaOS)3/2ClOH3N:NCSH4SONA-2 with 2-chloro-6-quinoxalinecarbonyl chloride and salting with Na2SO4.
18014-03-2P
RL: IMF (Industrial manufacture); PRSP (Preparation)

RR: INF (Industrial manufacture); PREP (Preparation)
(preparation of)
16014-03-2 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[4-(2,3-dichloro-N-ethyl-6quinoxalinecarboxamido)-3-sulfoanilino]-9,10-dihydro-9,10-dioxo-, disodium
salt (SCI) (CA INDEX NAME)

L13 ANSWER 163 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
1967:86597 CAPLUS
1NYENTOR(S):
Reactive ato dyes
Siegel, Régar; Samme, Klaus
PATENT ASSIGNEE(S):
SOURCE:
COPYRIGHT TYPE.

DOCUMENT TYPE: Patent German LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. DE 1233519

Azo dyes containing a 6-substituted-2,3-dichloroquinoxaline group (Q) and useful for dyeing cellulose, wool, polyamide, and polyeater fibers were prepared Thua, 34.7 parts 2,4.8-HANC10H5 (SONMs) 2 was diszotized and coupled with 10.7 parts 3-MecGH4NH2, the aminozo dye salted, filtered, washed, dissolved at pH 7 in 700 parts H20 by adding NaOH, attred at pH 7-7.5 with a solution of 25 parts (NNO in CSH6, salted, and filtered to give a feat reddish yellow dye for cotton. Similarly, the following dyes were prepared (reactants and shade on fiber given): QNH2 → 1-(3,5-dichloro-4-sulfophnyl)-3-methyl-5-pyrazolone, reddish-yellow on cellulose; QNH2 → PNOH, CHACH2OH12, orange on polyester and polyamide fibers; QNH2 → PNOH, Teddish yellow on polyamide fibers; 2-HO3SCGH4NH2 → PNOH, Teddish yellow on polyamide fibers; 2-HO3SCGH4NH2 → 1,3,6,8-HO(HO3S)2C10H4NHCOCH2NMeCOQ, bluish red on cellulose, wool, and

(preparation of)
1345-25-6 CAPLUS
2-Phenazinecarboxamide, N-[2-(dimethylamino)ethyl]-, 5,10-dioxide (SCI, 9CI) (CA INDEX NAME)

13458-27-0 CAPLUS
2-Phenazinecarboxamide, N-[3-(dimethylamino)-2-methylpropyl]-,
5,10-dioxide (SCI, SCI) (CA INDEX NAME)

13458-29-2 CAPLUS 2-Phenarinecarboxamide, N-[2-{1-pyrrolidinyl}ethyl}-, 5,10-dioxide (&CI, 9CI) (CA INDEX NAME)

13458-30-5 CAPLUS 2-Phenazinecarboxamide, N-[3-(dimethylamino)propyl]-, 5,10-dioxide (SCI, 9CI) (CA INDEX NAME)

polyamide fibers; 1,2,3,6,8,7-HO(QN:N) (HO3S)2(H2N)ClGH2N:NC6H3(SO3H)NH2-2,4, QNCO, greenish blue on celluloae.
14573-57-07
RE: INF (Industrial manufacture); PREP (Preparation)
(preparation of)
14573-57-0 CAPLUS
2,7-Naphthalenediaulfonic acid, 5-[2-(2,3-dichloro-N-methyl-6-quinoxalinecarboxamido) acetamido)-4-hydroxy-3-[(o-sulfophenyl)azo]- (7CI, sCI) (CA INDEX NAME)

L13 ANSWER 164 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
156:37957
66:37957
TITLE:
Phenaxine derivativea
PATENT ASSIGNBE(8):
Neth. Appl., 10 pp.
CODEN: NAXXAN
PAECAL

DOCUMENT TYPE: LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. APPLICATION NO. NL 1966-3503 FR DATE NL 6603503 FR 1462194 FR 89671 GB 1068985 US 3455926 PRIORITY APPLN. INFO.: 19660318 19650325 19660204 19690715

For diagram(s), see printed CA Issue.
The title compds. (I,X = NH2 or substituted-amino or N-heterocyclic modety) are prepared by reaction of phenazine-2-carboxylic acid 5,10-dioxide (II) with N,N'-carbonyldimidazole (III) to give I (X = 2-imidazolyl), which then is treated with an NH2 derivative Thus, to 17.2 g. III (68%) in 500 cc. dry HCONMe2 (IV) 12 g. II is added, the mixture kept 24 hrs. at normal temperature, 21 g. 1-methylpiperazine added, and after 4 hrs. the our

normal temperature, 11 g. 1-metry/piperazine school, and alter a first cooled to 5° to obtain 11.4 g. I (X = 4-methyl-1-piperazinyl), m. 209-10°. Similarly the following I were obtained [X and (m.p.) given]: X = 2-dimethylaminosthylamino (180°); 4-ethyl-1-piperazinyl (160-2°); 3-dimethylamino-2-propylamino (164-6°); 4-benzyl-1-piperazinyl (162°); (2-(1-pyrrolidinyl)ethylamino] (184-6°); 3-(dimethylamino)-propylamino (161-2°); [2-(4-methyl-1-piperazinyl)ethylamino] (170-2°). The compds. and their salts or quaternary N derive: are anticancer agents. 13458-25-8P 13458-27-0P 13458-23-2P 14559-63-8P RL: SPN (Synthetic preparation); PREP (Preparation)

14559-63-8 CAPLUS
2-Phenazinear-Phozamide, N-{2-(4-methyl-1-piperazinyl}ethyl}-, 5,10-dioxide
(BCI, 901) (CA INDEX NAME)

LI3 ANSWER 165 OF 181

ACCESSION NUMBER: 1366:51469 CAPLUS

COCUMENT NUMBER: 64:51469

COURNIN NUMBER: 64:51469

ACTIONER REFERENCE NO.: 64:9566d-f

TITLE: Structure vs. reactivity in quinoxalinecarboxylic acids and esters

AUTHOR(S): Gum, Wilson F., Jr.; Joullie, Madeleine M.

CORPORATE SOURCE: Univ. of Pennsylvania, Philadelphia

Journal of Organic Chemistry (1965), 30(11), 3982-5

COURNI TYPE: Journal

LANGUAGE: English

AB In an attempt to establish a correlation between the calculated electron densities in an unperturbed quinoxaline nucleus and the reactivities of its derive., the pKA values of 2-, 2,3-, 6-, 5-, and 2,3-dimethyl-5- quinoxalinecarboxylic acids (I, II, III, IV, V) were measured. The carbonyl frequencies of the corresponding Me and Et esters (VI-XII) were determined by it spectroscopy and tabulated together with those of Et and Me pyta-zinecarboxylate. Good correlation seemed to exist between pKA values of the acids I-V and the electron d. calculated by Longuet-Higgins and Coulson (CA 41, 4978b) but only poor correlation with those reported by Basur and Bhattacharya (CA 52, 864 and by Pullman (CA 41, 197b). The split carbonyl bands observed for the esters VI-XII should be ascribed to conformational iscemeriam rather than to Permi resonance. Relative pKA values of I-V were predictable from electron densities of the unsubstituted quinoxaline ring even though the carboxylate anions formed during the determination must perturb the ring denaities. The effect is apparently small in relation to the perturbed electron d. caused by the ring M acome in the unsubstituted quinoxaline nucleus.

15 6524-72-7, 6-00-dinoxalinecarboxylic acid, ethyl ester

(apectrum of, reactivity and)

RN 6524-72-7, CADUnoxalinecarboxylic acid, ethyl ester

(Apectrum of, reactivity and)

ANSWER 166 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN 1965:489504 CAPLUS 63:89504

TITLE: Asaporphine dyes
INVENTOR(S): Asaporphine dyes
Molf, Walther; Schroeter, Rudolf
PATENT ASSIGNEE(S): Street, Rudolf
PATENT ASSIGNEE
DOCUMENT TYPE: Detent
DATENT INFORMATION:

PATENT INFORMATION:

KIND DATE 19650312 APPLICATION NO. DATE

PATENT NO. KIND DATE APPLICATION NO. DATE

FR 192152 19650312 FR 1964-970502 19640410

BE 646314 GB

BE 646314 GB

BE 1020304 GB

PRIORITY APPLA INFO.:

OF Todingram(s), see printed CA Issue.

AB Reactive dyes for cotton having the general formula I (M is Co or Cu, 0 = Co or SO2, and Pc is a phthalocyanine residue) are prepared by condensing a tri- or tetra-sulfonyl chloride of Cu- or CoPe with an appropriate amine to give a sono- or diantiled, which is then ecylated on the terminal Mint group(s). Thus, a neutral paste of CuPe(3-503C1)4 (III), prepared from 300 g. 956 CuPe, was mixed with a small amount of water and the volume brought to 2 1. It was treated with a solution of 216 g. 4,2-HZN HGO3S C6H3CH3NHMe (III), m. 240-5° (decomposition) (from 4,2-HZN HGO3S) C6H3CH3NHMe (III), m. 240-5° (decomposition) (from 4,2-HZN HGO3S) C6H3CH3NHMe (IV), MeNHZ, and H) and 80 ml. 37% HCl, then dropwise during 2 hrs. at 0-3° with 150 ml. pyridine; the temperature rose slowly to 20° and continued to rise while dilute NsON was added dropwise during 1-1.5 hrs. until the diamilide was formed in solution at 25-30° and pR 6.0; after removal of the pyridine at pH 9 by steam distillation, 300 g.

2.3-dichloroquinoxaline-6sulfonyl chloride was added with agitation at pH 8-9 (attained by dropwise addition of dilute NsON), the dye kept in solution by addition of water up to total and after clarification and adjustment of pH to 7.0, precipitated with

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2 D1-so3-

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31157-47-8 CAPLUS
Cobaltate(3-), [c-[[4-[[(2,3-dichloro-6-quinoxaliny1)carbony1][(2sulfophny1)methy1]amino|methy1]pheny1]amino|aulfony1}-29H,31Hphthalocyanine-C.c-disulfonato(5-)-N29,N30,N31,N32]-, trihydrogen (9CI)
(CA INDEX NAME)

edisulfonato(2-)]- (?) 31215-23-3, Copper, [trihydrogen [[2-]3-(2,3-dichloro-N-sethyl-6-quinoxalinecarboxamido)butyl]-5-sulfophenyl]sulfemoyl]phthalocyaninedisulfonato(2-)]- 31226-61-2, Cobalt, [trihydrogen [[p-[2-12,3-dichloro-N-(4-sulfobtyl)-6-quinoxalinecarboxamido]ethyl]phenyl]sulfamoyl]phthalocyaninedisulfonato(2-)]-

(preparation of)
31133-54-4 (APRUS
([3-{[[3-{[[3-4]chloro-6-quinoxalinyl)carbonyl]methylamin
o|methyll-4-sulfophenyllamino|aulfonyll-39H,31H-phthalocyanine-C,Cdisulfonato(5-)-N29, No. N31, N327, trihydrogen (9C1) (CA IMDEX NAME)

2 D1-803-

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31132-55-5 CAPLUS
Cuprate(3-), [C-[[[4-[[[4-([[4-(1]] 2-dichloro-6-quinoxalinyl)carbonyl]methyllamino]methyll-1-sulfophenyl]methyll-1-sulfophenyllamino]methyll-1-sulfophenyllaminojmethyll-2-ynlyndyndinoxac(C-disulfonato(5-)-N29,N30,N31,N32]-, trihydrogen (9CI) (CA INDEX NAME)

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2 D1-so3-

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●3 H*

31215-23-3 CAPLUS
Cobaltate(3-), [C-{[{4-{2-{{(2,3-dichloro-6-quinoxalinyl)carbonyl}}(1sulfopropyl)aminojethyl]phenyl|aminojeulfonyl]-298,31H-phthalocyanine-C.Cdisulfonato(5-)-N29,N30,N31,N32}-, trihydrogen (9CI) (CA INDEX NAME)

2 D1-so3-

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●3 H+

31215-23-3 CAPLUS
Cuprate(3-), [c-[[[2-[3-[[(2,3-dichloro-6-quinoxalinyl)carbonyl]methylamin olbutyl]-5-aulfophenyl]amino]aulfonyl]-29H,31H-phthalocyanine-C,C-diaulfonato(5-)-N29,N30,N31,N32]-, trihydrogen (9CI) (CA INDEX NAME)

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2 D1-so3-

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L13 ANSWER 167 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1965:455160 CAPLUS
DOCUMENT NUMBER: 63:55160
ORIGINAL REFERENCE NO.: 63:10100f-h,10101a-b 1965:455160 CAPLUS
63:55160
63:10100f-h,10101a-b
Reactive dyes
Rothman, Leonard A.
E. I. du Pont de Nemours & Co.
11 pp.
Patent
Unavailable TITLE: INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: LANGUAGE FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. PATENT NO. FR 1384789 19650108 PR 1963-955250 19631127

GB 1000527 GB

US 3323931 19660201 US 1962-240747 19621128

PRIORITY APPLN. INFO: US 1962-240747 19621128

GB FOO diagram(s), see printed CA Issue.

AB Compds. of the general formula I, where Pc is phthalocyanine, Rl is H,

2 [D1-603-]

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●3 H+

31216-61-2 CAPLUS
Cobaltate(3-), [C-{[{4-[2-[[(2,3-dichloro-6-quinoxaliny1)carbony1](4-sulfobuty1)amino]ethy1]pheny1]amino]ethy1]pheny1]amino]ethy1]-29K,31H-phthelocyanine-C,C-disulfonato(5-)-N29,N30,N31,N32]-, trihydrogen (9CI) (CA INDEX NAME)

alkyl, or aryl, A is a p-C6H4 or sulfophenylene, R2 is H or St. R3 is a heterocyclic residue, and a * b * c * 3 * 4, dys cotton turquoise. Thus, 23 parts CuBe was added to 160 parts CLSO3H at 5° keeping the temperature below 25°, the mixture stirred 15 min., heated during 1.5 hrs. to 135 ; 5°, stirred 3.5 hrs. at that temperature, then the green solution cooled to room temperature, poured into a mixture of ice and H2O at <5°, the precipitate filtered and washed with 1 HCl at 5°, to give CuPc(1:603H)x(3:602Cl)y(x * y = 3 * 4) (II). 2,4 * (H2N) 2C6H3503H (III) (22.5 parts) was added to II in 1200 parts ice * H2O, the pH adjusted to 5 with 10 N NaOH, and then to 9.9 with aqueous NH3, the mixture heated to 25:30° and stirred until the PH stayed at 8.8-9.2 without further addition of NH3 (about 15 hrs.), acidified with HCl, the precipitate filtered, washed with ous

ous
HCl to remove excess III, the filter cake mixed with 1500 parts H2O at
35-40°, the suspension adjusted to pH 7.2 with NaOH, 21 parts
2,3-dichloro-6-quinoxalinecarbonyl chloride (IV) added, the mixture stirred
8 hrs. at 35-40°, keeping the pH at 7-7.5 with NaOH, the turquoise
solution (filtered, NaCl carefully added to 18%, the precipitate filtered,

solution filtered, NaCl carefully added to 18%, the precipitate filtered, and with 20% aqueous NaCl, and vacuum-dried at 60° to give I [R = Rl = R2 = N. A = 4 = N. A = 1.3. C = 1.3.

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L13 ANSWER 168 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
ORIGINAL REFERENCE NO.: 63:19368
ORIGINAL REFERENCE NO.: 63:7145b-f
Chromium- and cobalt-containing azo dyes
PATENT ASSIGNEE(8):
SOURCE: 7. R. Geigy A.-G.
23 pp.
DOCUMENT TYPE: Patent Unavailable

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO. APPLICATION NO. PATENT NO. KIND DATE APPLICATION NO. DATE

NL 4411791 19650412 NL 1964-11791 19641009

PRIORITY APPLM. INFO.:

AB 2.5-HD (DAIN CHIN. NCH. - NUNCSH4502NH2-m (I) 2.2. from equimoler smis. dissociated 2.4-HM (0201)CSH40NH (II) and m-HANSOGICHANNHX: CHIN. and dissociated 2.4-HM (0201)CSH40NH (II) and o-HACHACONH-CSH4CI (IV). in HCONNIZ 300 treated at 80-5* with Co(OAC) 2.15 parts and salted with Nacl gave a dark powder which dyes wool yellowish doi: 10.8 and III 12.7 parts gave a more grayish dye. Similar Co complex dyes were obtained from I 22 with IV -+ a-ACCHZOONHCSH4CI 21.2, with IV -+ ACCHZOONHCH 19.5, and with II -+ V 1.8. parts. 2.5-HO(OZN).

CCHIN. NCDh. NHHCSHHSOZNHCHZCHZOH-B 24.2 and 1.2-(2.4-HO (OZN)CSH3N.N)CIOHSON 15.5 in HCONNEZ 300 treated at 80-5* with Co(OAC) 2.15 parts gave a dark powder; vicioHenniZ -- 2-CIOHFOH in KONNEZ 333 treated with 0-CHZCCGH4N: NCDh. NNHDCH NNHDN (VII) 17.2 and heated with Na2CO3 15 parts at 100-5* gave a dark powder; it dyes gray shades. The same dye was also obtained by treating VI 22.3 parts in RTOH 800 vols. and H2O 150 parts with VII 17.2 parts and 10N NoH 15 vols. and heating at 60-5*. Similar dyes were prepared from VI 22.3 with 2.5-HAN(PhNHSOZ)-CSH3COZH -> VIII 25.7, or with 4.3-HO(H2N)-CSH3COZH -> VIII 25.7, or with 4.3-HO(H2N KIND DATE DATE

substituted a-triazinyl groups were prepared p-HANCGCH2NHMe (9.85 parts) diszotized and coupled with 12 parts 8,4,6,1-HO(HO35)2C10H4NHCOCH2SO3H and the mixture treated at 15° with 2,3-dichloroquinoxaline-6-carbonyl chloride (11 9 and Na2CO3 7.7 in H2O 40 parts yielded II. 4,2-H2N(HO35)2GH3CH3NHMe (12.5 parts) diszotized and coupled with 21 parts 8,3,6,1-H0(HO35)2GH3CH3NHMe (12.5 parts) diszotized and coupled with 12.5 parts I gave a dye which dyed cotton and regenerated cellulose fabrics brilliant bluish red shades of good wet- and lightfastness. 3,4 H3N(HO35)CGH3CH3NHME (III) (12.5 parts) diszotized and coupled with 21.8 parts 5,2,1,7-N(HANCONH)C10H4(SOAH)2, and the mixture treated at 40-5° with I gave an orange-red dye. III (12.5 parts) diszotized and coupled with 19.9 parts 9,7-2HO(HO35)CGHSNHME (IV), and the mixture condensed with 10.7 parts cyanuric chloride in 90 parts M32CO gave a brilliant orange dye. III (12.5 parts) diszotized and coupled with 12 parts 4,1- HO35COH6CH and then condensed with 13 parts 2,4-dichloropyrimidine-6-carbonyl chloride gave a brilliant scarlet dye. III(11.5 parts) diszotized and coupled with 12.12 parts | Jack10H3D(SCH3CH2 NHME (12.5 parts) diszotized and coupled with 12.7 parts p-202HCH3HM2 8,1,6,1-HO(HO35)2CIOH4NH2 and then condensed with 15.2 parts I gave a blue dye. III (12.5 parts) diszotized and coupled with 15.7 parts p-202HCH4HM2 8,1,6,1-HO(HO35)2CIOH4NH2 and then condensed wiccessively with 10.7 parts cyanuric chloride and olparts m-HO35CGH4NH2 gave a brilliant orange dye.

2752-29-6 CAPLUS
1,7-Maphthalenediaulfonic acid, 6-[[p-[(2,3-dichloro-N-methyl-6-quinoxalinecarboxamido)methyl]phenyl]azo]-5-hydroxy-4-(2-sulfoacetamido)-, trisodium salt (SCI) (CA INDEX NAMS)

2.4-HO(PhSO2)C6H3N:CHC6H4OH-O 17.7 in HCONN2300 with Co(OAc)2 15 parts at 80-5° gave a dark powder, olive-green on wool. Similar dyes were obtained from 1X 22 with 2.4-HO(PhN:N)C6H3CH:NCCH3-(OH)NO2-2.5 18.1 or with 2.4-HO(PhN:N)C6H3CH:NCCH3-(OH)NO2-2.5 18.1 or with 2.4-HO(PhN:N)C6H3CH:NCCH3-(OH)NCGH2-2.5 19.4 parts.
3.2,5-C1.400(OH)NCGH2N:NCCh1:NCH6-H802NM2-2 21.7 and IV →
3-methyl-1-phenyl-3-pyraxolone (X) 19.3 with Co(OAc)2 15 parts gave a dark powder, brown-olive on wool. Similar Co complex dyes were prepared from mixta. of 5,2-C1(HO)C6H3N:NCCh1:NHCK6H802NM2-m (XI) 21.5 and 1,2-[2,5-HO(C3H0C6H3N:N)CCh1:NHCK6H802NM2-m (XI) 21.5 and 2.5,4-Ho(Ex502)(O2N)C6H2N:NCCh1:NHCK6H802NMHe-p 27.3 and of 2.5,4-HO(N)(PhN:N)C6H2NN-NCDh1:NHCK6H802NMHe-p 27.3 and of 3.2,4-HO(N)(PhN:N)C6H2NN-NCDh1:NHCK6H802NMHe-p 27.3 and of 3.2 and 111 21.2 parts (olive-green). I 22 and 2,5-HO(M502)C6H3NN2 → 2,7-HOCLOH5NNHC-20 with Cr-(OAc)216 and Na2CO3 15 parts in HCONN2333 vols. gave an olive-gray dye.
4445-61.8 0.7-Naphthalenedisulfonic acid, 4-amino-6-[(α-(2,3-dichloro-N-methyl-6-quinoxalinecarboxamido)-4-sulfo-m-tolyl]azo]-5-hydroxy-3-[(p-nitrophenyl)azo]- (7CI, SCI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

CAPLUS COPYRIGHT 2006 ACS on STN 1965:439567 CAPLUS 63:99567 63:7144g-h,7145a-b Monoazo dyee Parbenfabriken Bayer A.-G. L13 ANSWER 169 OF 181 ACCESSION NUMBER: DOCUMENT NUMBER: ORIGINAL REFERENCE NO.: TITLE: PATENT ASSIGNEE(S): 21 pp. Patent Unavailable SOURCE: SOURCE: DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. PATENT NO. KIND DATE DATE NL 6410555 196409
PRIORITY APPLN. INFO.: DB 196309
AB AZO dyes contg 2,3-dichloro-6-quinoxalinyl, dior trichloropyrimidyl,

L13 ANSWER 170 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1965:439122 CAPLUS COCUMENT NUMBER: 63:39122 CAPLUS 63:7011c-f

63:7011c-f
Some substituted 1,2,3,4-tetrahydroquinoxalines and
Hofmann degradation of a quaternary ammonium hydroxide
derived from N,N'-dimethyltetrahydroquinoxaline
Elina, A. S.; Musatova, I. S.
S. Ordshonkidze All-Union Chem.-Pharm. Research
Inst., Moscow
Khimiya Geterotsiklicheskikh Soedinenii (1965), (2),

AUTHOR(S): CORPORATE SOURCE:

SOURCE:

CODEN: KGSSAQ; ISSN: 0132-6244 Journal

291-5
CODEN: KOSSAQ; ISSN: 0132-6244
DOCUMENT TYPE: Journal
LANGUAGE: Russian
AB N.N.N.*Trimethyl-1,2,3,4-tetrahydroquinoxalinium iodide (I) undergoes a degradation in alkaline solution with the formation of methylated o-phenylenediamine deriva. N-Acetyl-1,2,3,4-tetrahydroquinoxaline and PhCHGIC yielded a mixture of products: N-acetyl-1,2,3,4-tetrahydroquinoxaline, m. 60.5-2.5°; N-benzyl-1,2,3,4-tetrahydroquinoxaline, m. 60.5-2.5°; N-benzyl-1,2,3,4-tetrahydroquinoxaline, m. 60.5-2.5°; N-benzyl-1,2,3,4-tetrahydroquinoxaline, m. 92-6°; and N.N'-dibenzyl-1,2,3,4-tetrahydroquinoxaline, m. 92-6°; and N.N'-dibenzyl-1,2,3,4-tetrahydroquinoxaline, m. 92-6°; and N.N'-dibenzyl-1,2,3,4-tetrahydroquinoxaline, m. 101-000 (Red and Notherhyl-1,2,3,4-tetrahydroquinoxaline) iodide m. 175-6°. I and 40% solution of NaOH refluxed 6 hrs. gave N.N.N'-trimethyl-o-phenylenediamine (III), hi 62-4° (picrate m. 112-13°; HCl salt m. 165-7°) and N.N'-dimethyl-1,2,3,4-tetrahydroquinoxaline, bi 92-4°; picrate m. 122-4°.
Distillation of I gave III. Mei and III gave
2-methylaminophenyltrimethylammonium iodide, m. 219-20° (decomposition), the product of the reaction of N.N.N'.N'tetramethyl-o-phenylenediamine with MeI.
II 2427-91-0, 6,7-Quinoxalinedicarboxylic acid, 5,8-dihydroxy-, diethyl ester (preparation of)
RN 2427-91-0 CARUUS
CN 6,7-Quinoxalinedicarboxylic acid, 5,8-dihydroxy-, diethyl ester (7CI, 8CI) (CA INDEX NAME)

L13 ANSWER 171 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1965:439121 CAPLUS
CONCENT NUMBER: 63:39121
CRIDINAL REFREENCE NO: 63:39121
AUTHOR(S): Chelating reagents containing nitrogen heterocycles.
IV. Syntheses of 5.8-dihydroxyquinoxaline derivatives
CORORATE SOURCE: Nippon Kegaku Zesehi (1965), 86(4), 435-7
CODEN: NRPEAZ: 125N: 0369-5387
DOCUMENT TYPE: Journal
LANGUAGE: Japanese
Cf. CA 63, 4295c. 2,3-Dichloro-5,8-dimethoxyquinoxaline (I) (1 mole) and
2.4 g. atom Ns in appropriate alc. was heated 3 hrs. to give the following
2,3-dislkoxy-5,8-dimethoxyquinoxalines (alkyl in alkoxyl, m.p., and %
yield given): Et. 172*, 73; EXC(CM2)2, 114*, -- Heating I
with 404 aqueous Ne2NNI at 120-30* for 3 hrs. gave 2,3bis(dimethylamino)-5,8-dimethoxyquinoxaline, m. 16-17*. Using 1
mole 2,3-dichloro-5,8-dihydroxyquinoxaline (alkyl, m.p. and m.p.
of acetate given): Et. 148* 168*; EXC(CM3)2, 65*,
78*. Similarly, 2,3-bis(dimethylamino)-5,8-dimydroxyquinoxaline, (alkyl, m.p. and m.p.
of acetate given): Et. 148* 168*; EXC(CM3)2, 65*,
78*. Similarly, 2,3-bis(dimethylamino)-5,8-dimydroxyquinoxaline, m.
151-2*, and 2,3-bis(dimethylamino)-5,8-dimydroxyquinoxaline, m.
151-2*, and 2,3-bis(dimethylamino)-5,8-dimydroxyquinoxaline, m.
2, di-Et pyrazine-2,3-dicarboxylate, b) 165*, n200* 1.5559,
and 1.0 g. powdered Ns in xylene was heated in 1n at 110* to give
2,7 g. di-Et 5,8-dihydroxyquinoxaline-6,7-dicarboxylate, m. 145*,
dicarboxylate, m. 183-4*, was obtained in 30* yield. Similar
treatment of di-Et 2,3-dimydroxyquinoxaline-6,7-dicarboxylate, with II failed to
give quinoxaline derivative but gave di-Et cyclohexane-2,5-dione-1,4dicarboxylate, e. 2452-36-0, 6,7-Quinoxalinedicarboxylic acid.

give quinoxaline derivative but gave di-Et cyclohexane-2,5-dione-1,4-dicaxboxylate.

2427-91-0, 6,7-Quinoxalinedicarboxylic acid, 5,8-dihydroxy-,
diethyl ester 2452-36-0, 6,7-Quinoxalinedicarboxylic acid,
5,8-dihydroxy-, diethyl ester, diacetate (ester)
(preparation of)
2427-91-0 CAPLUS
6,7-Quinoxalinedicarboxylic acid, 5,8-dihydroxy-, diethyl ester (7CI, 8CI)
(CA INDEX NAME)

2452-36-0 CAPLUS 6,7-Quinoxalinedicarboxylic acid, 5,8-bis(acetyloxy)-, diethyl ester (9CI)

1910-93-6 CAPLUS
2,3-Phenazinedicarboxylic acid, 1,4-dihydroxy-7,8-dimethoxy-, dibutyl ester, diacetate (ester) (SCI) (CA INDEX NAME)

1910-94-7 CAPLUS 2,3-Phenazinedicarboxylic scid, 7,8-dibutoxy-1,4-dihydroxy-, diethyl ester (7CI, SCI) (CA INDEX NAME)

1910-95-8 CAPLUS
2,3-Phenazinedicarboxylic acid, 1,4-bis(acetyloxy)-7,8-dibutoxy-, diethyl ester (9C1) (CA INDEX NAME)

1983-91-1 CAPLUS 2,3-Phenazinedicarboxylic acid, 1,4-dihydroxy-7,8-dimethoxy-, dibutyl ester (7C1, 8C1) (CA INDEX NAME)

(CA INDEX NAME)

L13 ANSWER 172 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1965:424143 CAPLUS DOCUMENT NUMBER: 63:24143 CAPLUS ORIGINAL REFERENCE NO.: 63:4295c-f

es:+4:350-12
Chelating reagents containing nitrogen heterocycles.

I. Syntheses of 1,4-dihydroxyphenazine derivatives
Oguchi, Shoichi
Tokyo Gakugei Univ.
Nippon Kagaku Zasshi (1965), 86(2), 246-9
CODEN: NPKZAZ; ISSN: 0369-5387

AUTHOR(S): CORPORATE SOURCE: SOURCE:

DOCUMENT TYPE: LANGUAGE:

CODEN: NPKZAZ; ISSN: 0369-5387

JUMENT TYPE: Journal

Jumenal

Jumenal

Jumenal

Jumenal

Jumenal

Jumenal

Jumenal

Jumenal

The optimal conditions for condensation of di-Et quinoxaline-2,3dicarboxylate (1) with (CH2COZEL)2 (II) were sought. Using NaOEt, NaNH2,
PhlCNa, and NaH as base, 124 di-Et cyclohexane-2,5-dione-1,4dicarboxylate, quinoxaline-2,3-dicarboxamide, tar, and 35t di-Et
1,4-dihydroxyphenazine-2,3-dicarboxylate (III) were obtained, resp. The
best yield (511) of III was obtained when 1:1:2 molar ratio of I, II, and
Na or K was used. 4,5-(CNN)2CER(208)2-1,2 (9.2.9., 25 g. Sn. 130 ml.
concentrated HCl, and 60 ml. BCDH was heated to give 4,5-(RNN)2CENI2(OB)2-1,2

(IV) m. apprx.95° (decomposition); di-Ac derivative m. 162°.

Treatment of IV with [HO2CCON] 2] gave 6,7-dibutoxyquinoxaline-2,3dicarboxylic acid, m. 148-50° (decomposit) di-Et ceiter b)

124°, m. 95-6°. Similarly, 6,7-dimethoxyquinoxaline-2,3dicarboxylic acid, m. 128-2 prepared VIII. and Na in 1:1:2 molar
ratio was treated as above to give 474 di-Et 1,-dihydroxy-7,8dimethoxyphenazine-2,3-dicarboxylate, m. 256-7°. disectate, m.
216-19°. Similarly, di-Bu 1,4-dihydroxy-7,8-dimethoxyphenazine-2,3-dicarboxylate, m. 233-4° (diacetate m. 165-6°), and di-Et
1,4-dihydroxy-7,8-dibutoxyphenazine-2,3-dicarboxylate, m. 181-2°
(diacetate, m. 26-7°), were prepared Similar treatment of I with
(CH2CN)2 gave 744 2,3-dicyno-1,4-dihydroxy-7,8-dimethoxy-1

1770-41-8, 2,3-Phenazinedicarboxylic acid, 1,4-dihydroxy-7,8dimethoxy-, dicthyl ester 1910-93-6, 2,3-Phenazinedicarboxylic
acid, 1,4-dihydroxy-7,8-dimethoxy-, dibutyl ester, diacetate (ester)
1910-94-7, 2,3-Phenazinedicarboxylic acid, 1,4-dihydroxy-1,4-dihydroxy-7,8-dimethoxy-, diethyl ester, diacetate (ester)
1910-94-7, 2,3-Phenazinedicarboxylic
acid, 1,4-dihydroxy-7,8-dimethoxy-, diethyl ester, diacetate (ester)
1910-94-7, 2,3-Phenazinedicarboxylic
acid, 1,4-dihydroxy-7,8-dimethoxy-, diethyl ester, diacetate (ester)
1910-94-7, 2,3-Phenazinedicarboxylic
acid, 1,4-dihydroxy-7,8-dimethoxy-1

(prepar

3684-53-5 CAPLUS 1,2,3,4-Phenazinetetrol, 7,8-dimethoxy-, tetrascetate (ester) (SCI) (CA INDEX NAME)

L13 ANSWER 173 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1963:448886 CAPLUS

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.:

1963:448886 CAPLUS 59:48886 59:8910e-h Water-soluble anthraquinone reactive dyes Singer, Josef; Schwechten, Heinz N. Farbenfabriken Bayer A.-G. 18 pp. Patent Unavailable TITLE: INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: LANGUAGE:

PATENT INFORMATION:

APPLICATION NO.
BE
GB PATENT NO. KIND DATE

PATENT NO. KIND DATE APPLICATION NO. DATE

BE 622859 19630115 BE 19630125 BE 1964025 1964025 BE 622859 1964051 BE 1964051

RN 104242-65-1 CAPLUS
CN 2-Anthracenesulfonic acid, 1-emino-4-[α-(2,3-dichloro-N-methyl-6-quinoxalinecerboxamido)-p-toluidino)-9,10-dihydro-9,10-dioxo-, sodium selt (7C1 (CA 170EX NAME))

104601-65-2 CAPLUS
2,6-Anthracenedisulfonic acid, 1-amino-4-[u-(2,3-dichloro-N-methyl-6-quinoxalinearboxamido]-p-toluidino]-9,10-dihydro-9,10-dioxo-, disodium salt (7CI) (CA INDEX NAME)

PAGE 1-A

●2 Na

104601-66-3 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-{[4-{[[(2,3-dichloro-6-quinoxaliny])carbony]]methylamino]methyl]-1-sulfophenyl]amino]-9,10-dihydro-9,10-dioxo-, disodium salt (9CI) (CA INDEX NAME)

105232-45-9 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[3-[3-[2,3-dichloro-N-methyl-6-quinoxalinacrboxamido)-5-methylhexyl]sulfoanilino]-9,10-dihydro-9,10-dioxo-, disodium salt (7CI) (CA INDEX NAME)

D1-803H

●2 Na

106303-94-0 CAPLUS
2-Anthracenesul(onic acid, 1-amino-4-[4-[3-(2,3-dichloro-N-methyl-6-quinoxalinecarboxamido)propyl]-3-sulfoanilino]-9,10-dihydro-9,10-dioxo-,disodium salt (7CI) (CA INDEX NAME)

106337-79-5 CAPLUS
2-Anthracenseulfonic scid, 1-smino-4-[4-{[4-(2,3-dichloro-N-methyl-6-quinoxalineerboxamido]butyl]thio]-1-sulfosnilino]-9,10-dihydro-9,10-dioxo, disodium salt (7CI) (CA INDEX NAME)

PAGE 2-A

106381-98-0 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[4-[4-[4-[2,3-dichloro-N-methyl-6-quinoxalinoer>boxamido]butoxy]-3-sulfoanilino]-9,10-dihydro-9,10-dioxo-,disodium salt (7CI) (CA INDEX NAME)

⊕2 Na

107062-63-5 CAPLUS
2-Anthracenesulfonic acid, 1-amino-4-[3-[3-(2,3-dichloro-N-methyl-6-quinoxalinecerboxamido]butyl]sulfoanilino]-9,10-dihydro-9,10-dioxo-,disodium salt (7CI) (CA INDEX NAME)

D1- SO3H

●2 Na

L13 ANSWER 174 OF 181 CAPLUS COPYRIGHT 2006 ACS on 8TN
ACCESSION NUMBER: 1963:429002 CAPLUS
DOCUMENT NUMBER: 59:39002
ORIGINAL REFERENCE NO.: 59:52996-f,5300a-c
TITLE: OLINOXALINE dyes
INVENTOR(S): 59:52996-f,5300a-c
OLINOXALINE dyes
INVENTOR(S): 50:5296-f,5300a-c
OLINOXALINE dyes
INVENTOR(S): 50:5296-f,5300a-c
OLINOXALINE dyes
INVENTOR(S): 59:52996-f,5300a-c
OLINOXALINE dyes
INVENTOR(S): 59:52996-f,5300a-c
OLINOXALINE dyes
INVENTOR displayed
INV LANGUAGE: PATENT INFORMATION:

PAGE 2-A

●2 Na

106381-99-1 CAPLUS
2,6-Anthracenedisulfonic acid, 1-amino-4-[m-[[4-(2,3-dichloro-N-methyl-6-quinoxalinecarboxamido]butyl]sulfonyl]anilino]-9,10-dihydro-9,10-dioxo-, disodium salt (7C1) (CA INDEX NAME)

PATENT NO. KIND DATE APPLICATION NO. DATE

PRIORITY APPLN. INFO.:

AB Condensation compde. of 3- or 2-chloro- and 2.3-dichloroquinoxaline derivs. with azo, anthraquinone, or phthalocyanine dyes are H2O-soluble and suitable for dyeing or printing cellulosic and other materials. Thus, 2.4.6-H2N(Nox3)52(10HS [1] 34.7 was diszotized and coupled with 3-H2NC6H4Me [II] 10.7 parts. The product in equeous NaOH was mixed and stirred with 26.5 parts 2.3 dichloro-6-quinoxalinecarbonyl chloride [III] (m. 116*, bo.05 144*) in 100 parts C6H6, heated to 15

40*, neutralized with Na2CO3, and 80 parts NaCl added. The dye was filtered and dried at 40-50*. Printed on cellulose, it gave a reddish yellow color, fast to washing and light. Similarly, dyes were prepared (reactants and color of dye on cellulose, it gave a reddish yellow. 21-00.3-dichloro-6-isocyanacoquinoxaline. meddish yellow. 21-01. 3-dichloro-6-isocyanacoquinoxaline. meddish yellow. 2(1), reddish yellow. 21-02.7-dichloro-6-isocyanacoquinoxaline. meddish yellow. 2(1), reddish yellow. 21-02.7-dichloro-6-isocyanacoquinoxaline. meddish yellow. 2(1), reddish yellow. 21-02.7-dichloro-6-isocyanacoquinoxaline. meddish yellow. 2(1), reddish yellow. 2(1), r

L13 ANSWER 175 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1961:17956 CAPLUS
DOCUMENT NUMBER: 55:17956
ORIGINAL REFERENCE MO: 55:13796 didthyl 1,4-dihydroxy-5,8-dialkoxyphenazine-2,3-dicarboxylates
AUTHOR(S): Kewai, Sinichi; Torigoe, Masao; Fujiki, Shun; Shibata, Kiyoko; Oraki, Atauko; Sakakibara, Yoshiaki; Oguchi, Shoshichi
CORPORATE SOURCE: Nippon Kagaku Zasshi (1959), 80, 788-91
CODEN: NNEXAZ; ISSN: 0369-5387
DOCUMENT TYPE: Journal LANGUAGE: Unavailable

CODR: NPKZAZ; ISEN: 0369-5387

UNENT TYPE: Journal

UNADE: Unavailable

Hydroquinone (I) (210 g.), 565 g. PrBr, 2.1 l. dry acetone, and 750 g.

KZCO3 refluxed 50 hrs., the solvent removed, 4 l. H2O added, and the mixture

extracted with ether gave 271 g. hydroquinone dipropyl ether (III,).

50.5° (on evaporation of ether). Nitration (with HNO3 in AcOH) of 24 g. hydroquinone distropyl ether (III) yielded 28.6 g. crude nitro compound

(m. 176-7°); mechanical separation of the crystals gave the dinitro compound (V) of III, m. 141.5°. Similarly, II gave the dinitro compound (V) of III, m. 141.5°. Similarly, II gave the dinitro compound (V) of III, m. 141.5°. Similarly, II gave the dinitro compound (V) of III, m. 141.5°. Similarly, II gave the dinitro compound (V) of III, m. 141.5°. Similarly, II gave the dinitro compound (V) of III, m. 141.5°. Similarly, II gave the dinitro compound (V) of III, m. 141.5°. Similarly, II gave the dinitro compound (V) of III, m. 141.5°. Similarly, II gave the dinitro compound (V) of III, m. 141.5°. Similarly, II gave the dinitro compound (V) of III, m. 141.5°. Similarly, III gave the dinitro compound (V) of III, m. 141.5°. VI was condensed (VI), m. 139.46°. VI was condensed (VII), m. 139.46°. VI was condensed with dihydroxytartaria caid-Na to give 318 3. distehboxyquinoxaline-2, dicarboxylic acid (IX), m. 191.5°. according to the procedure of Adachi (CA Si, 17936b). IX condensed with dihydroxytartaria caid gave 92% 5.8-dipropoxyquinoxaline-2,3-dicarboxylia to the diethyl ester (XII), m. 157° (decomposition), which was converted to the diethyl ester (XII), m. 68.5-69°, by the usual method. Diethyl quinoxaline-2,3-dicarboxylate, m. 150°. According to the procedure of 68.5-69°, by the usual method. Diethyl quinoxaline-2.3-dicarboxylate, m. 150°. according to the procedure of 68.5-69°, by the usual method biethyl succinate, 15 cc. xylene, and 1.5 g. finely powdered No heated at 150-160° (5 hrs.) yielded diethyl 1,4-dihydroxy-5,8-dicarboxylate, m. 157°, and diethyl 1,4-dihydroxy-5,8-dicar DOCUMENT TYPE: LANGUAGE: AB Hydroquine

(preparation of)

cc. each of glaciel AcOH and RtOH treated in S hrs. with 70 g. PhNO gave 70 g. 5,2-Cl(AcNH)CSH3N:NPh (XXXV), yellow needles, m. 180° (from alc.); similarly was prepared 244 2.5-AcNH (Meo)CSH3N:NPh (XXXVI), orange blades, m. 174-5° (from EtOAC). XXXV (68 g.) and 700 cc. 5% alc. KOH refluxed 5 hrs. and the mixture poured into 2 l. H20 gave 70% 2-H2N analog, red needles, m. 113° (from petr. ether); similarly was prepared 69% 2.5-H2N(Meo)CSH3N:NPh, red laminase, m. 38.0-9.5° (from petr. ether). 4,2-(H2N)2CSH3Cl (28 g.), 23 g. 1,2-cyclohexanedione, and 240 cc. 10% aqueous AcOH heated 1 hr. at 98-100°, the mixture cooled, made alkaline with aqueous NaOH, and the precipitated solid filtered off, ed with

petr. ether). 4,2-(HZN)ZCGHZCI (28 g.), 23 g. 1.2-cyclohexanedione, and 240 cc. 104 aqueous ACOK heated 1 hr. at 98-100*, the mixture cooled, made alkaline with aqueous NaOK, and the precipitated solid filtered off, hed with HZO, dried, and extracted with petr. ether left an insol. compound (XXXVI), CLBHISMACI2, pale brown prisms, m. 274-6* (from ECOH) [presumably either 1,2-(2-smino-x-chlorophenylimino)cyclohexane (XXXVII) or R.C.C.(R), (CH2)4 (R = x-chloro-o-phenylendisimio) (XXXVIII); the petr. ether exts. gave 11.5 g. 7-chloro-1,2 3,4-tetrahydrophenazime (XXXIX), m. 94*. 2-HZNCSHANZPH (10 g.), 100 cc. cyclohexanone (XL), and 0.5 cc. concentrated HCl refluxed 2 hrs. and distilled until the temperature of mixture rose to 175*, the residue dissolved in Et2O, extracted with N HCl, and the HCl exts. made alkaline gave 6.9 g. 1,2,3.4-tetrahydrophenazine, b7 125-56*, m. 92-3* (from petr. ether); similarly, 4-15,2-Mc(EXN)(SIN)(SIS)(SIS) and XL gave 1,2,3.4-tetrahydro-7-methylphenazine (XLII), b5.5 163-4*, m. 80-2* (from petr. ether). Also prepared were 2,3-cyclopentenoquinoxaline (XLIII), b4 110-50*, m. 99-100*, 6-methyl-2,3-cyclopentenoquinoxaline (XLIII), b4 110-50*, m. 99-100*, 6-methyl-2,3-cyclopentenoquinoxaline (XLIII), b5 m. 103-4*, m. 81*. XLII (4.25 g.) in 75 cc. Et2O and 170 cc. 0.33M o-ROZCGM4COBH in Et2O kept in the dark 3 days, the N-oxide pth halate filtered off, washed with 8t2O, dissolved in H2O, the H2O solution adjusted to pH 7 with NaCOO, extracted with CRCl3, and the CRCl3 exts. dried and concentrated gave the 1,4-dioxide, green-yellow leaflets, m. 118-6* (from ECOH), XLIV, m. 204-6* (from ECOH), XLIV person end external call the sea unstable to VIITA. VIIIA by the above procedures gave 1,4-dioxides of the following compdex: XLI, yellow microcrystals. m. 186-8* (from ECOH); XLIV, m. 204-6* (from ECOH); XLIV, brown needles, m. 173-3* (from CGH6); and bornyleno(2*,3*,2,3)quinoxaline (resperation of) 10933-89-1, 6*Ouinoxalinecarboxamide, 2,3-dimethyl-N-(3-piperidinopropyl)-, 1,4-dioxide (6CI) (CA I

L13 ANSWER 177 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1957:1858 CAPLUS
DOCUMENT NUMBER: 51:1858
ORIGINAL REFERENCE NO.: 51:432e-1,433a-f Quinoxaline N-oxides. V. Purther bz-substituted derivatives 110081-11-3 CAPLUS
2,3-Phenazinedicarboxylic acid, 1,4-dihydroxy-6,9-dimethoxy-, diethyl ester (6C1) (CA IMDEX NAME)

113752-03-7 CAPLUS
2,3-Phenezinedicarboxylic acid, 6,9-diethoxy-1,4-dihydroxy-, diethyl ester (6CI) (CA INDEX NAME)

114399-30-3 CAPLUS
2,3-Phenaxinedicarboxylic acid, 1,4-dihydroxy-6,9-dipropoxy-, diethyl eater (ECI) (CA INDEX NAME)

ACCESSION NUMBER:

DOCUMENT NUMBER:

DOCUMENT NUMBER:

DOCUMENT NUMBER:

DITTLE:

Outhookaline N-oxides. VI. N-Oxides of 2,3-polymethylenequinoxalines

AUTHOR(g):

Landquist, Justus K.

TOPER JOURNAL JOURNA

AUTHOR(S): CORPORATE SOURCE: SOURCE:

Silk, J. A. Imperial Chem. Ltd., Manchester, UK Journal of the Chemical Society (1956) 2058-63 CODEN: JCSOA9; ISSN: 0368-1769

DOCUMENT TYPE:

DOCUMENT TYPE: JOUITMAI
LANGUAGE: Unavailable
AB 4.3-H2N(O2N)C6H3ON (from 4.3-AcNN(O2N)C6H3OAc and 6N HCl] (15 g.) in 150
cc. MeON hydrogenated over Raney Ni, filtered from the catalyst, the
filtrate concentrated to 75 cc. in vacuo, mixed with 10 g. anhydrous AcONA and

28.5
g. Glyoxal (XVIII) bisulfite in 140 cc. warm H2O, and the mixture heated 2.6
hrs. at 50° gave 7 g. 6-hydroxyquinoxaline (XIX), m. 252-4°
(from H2O); Ac2 in place of XVIII gave the 2.3-He2 derivative (XX) of XIX, m.
247-9° (from H2O). Na (0.46 g.) in 20 cc. EtOH treated with 2.92
g. XIX, then with 3.0 g. CLG12CO2Et, the mixture heated 2 hrs., cooled,
concentrated, poured into H2O, extracted with C6H6, and the C6H6 exts.
concentrated gave
0.9 g. 6-EtO2CCH2 derivative of XIX, m. 99-100° (from H2O),
5-Ethoxy-2,3-dismethylquinoxaline (5 g.) in 125 cc. C6H6 and 5 g. crushed
AlCl3 refluxed 16 hrs., the mixture cooled, decomposed with ice H2O, the C6H6
evaporated by air since the emulsion could not be broken, the solid (XXI)
c6H6,

the C6H6 exts. extracted with hot dilute aqueous NaOH, the NaOH exts.

the CGHS exts. extracted with hot dilute aqueous NaOH, the NaOH exts.

neutralized,
the solid (XXII) filtered off, and XXI and XXII recrystd. from H2O gave
0.89 g. 5-NO analog (XXIII), needles, m. 146-7°.
4.3-H2M(02H)(CGHSCO2Et, m. 140-2° [obtained in 70-85\$ yield from
4.3-AcMH-(O2H)(CGHSCO2Et) with (a) EtOH and 3\$ (volume/volume) H2504 or (b) with
EtOH-HCH], hydrogenated as above gave 30\$ 4.3-(ERM)ZCGHSCO2Et (XXIV),
needles, m. 112-14° (from dilute alc.). XXIV (4.5 g.) and (CGN)2
[from 6.1 g. sulfate (XXV) 40 cc. H2O, and BacO3] stirred vigorously 1
hr. at 60°, an equal portion of (CHO)2 added, and the stirring
continued 1 hr. gave 2.3 g. Et 6-quinoxalinecarboxylate (XXVI), m.
68-70° (from CGH6-cyclohexane, Al2O3); XXVI and VIIIa at room temperature
or at 50° gave an unidentified solid, m. sbout 340°. XXIV

(42 g.), 22 g. Ac2, and 500 cc. 33 EtOH refluxed 30 min. gave 48 g.
2.3-Me2 derivative (XXVII) of XXVI, feathery needles, m. 102-4°. XXVII
(5 g.) and 15 g. Et2N(CH2)2OH refluxed 16 hrs. and distilled gave 2.5 g.
EE2NGECH2 ester. m. 43-6° (by chromatography on Al2O3) in
CGH6-petr. ether). 4.3-H3N(O2H)CGHJAC (4.5 g.) hydrogenated in EtOH over
Pd-C, the mixture heated 1 hr. at 60° gave 1.4° g.
6-acctylquinoxaline (XXVIII), m. 106-6° (from 6 g.
XXVI), and the sixture heated 1 hr. at 60° gave 1.4° g.
6-acctylquinoxaline (XXVIII), m. 106-6° (from 6 g.
XXVI). insilarly was prepared the 2,3-Me3 derivative of XXVIII. m. 116-18° (from aqueous EtOH). Liklie (0.28 g.) in 50 cc. dry EtOH extend with 5 g. XXVII in
100 cc. dry EtOJ in 10 min., the mixture stirred 10 min., 2 cc. EtOAc added, the filtrate of the EtOJ leyer apparated, the aqueous extracted with EtO). Che mixture filtered, the EtOJ leyer apparated, the aqueous

layer extracted with Bt20, and the combined Bt20 exta. and solution dried and concentrated gave 0.7 g. 6-HoCN2 analog of XXVII, m. 113-14*.

gave 0.7 g. 6-HoCN2 analog of XXVII, m. 113-14*.

gave 0.7 g. 6-HoCN2 analog of XXVII, m. 113-14*.

show gave the 6-Ho analog (XXIX), yellow needles, m. 247-50*.

(decomposition) (from H20); 3 g. 6-hydroxy-2,3-dimethylquinoxaline and 75 cc. M VIIIA kept 17 hrs. at 60* gave 1 g. 1,4-dioxide (XXIXA), m. 249-50*. 5-Methoxy-2,3-dimethylquinoxaline 1,4-dioxide (XXIXA), m. 249-50*. 5-Methoxy-2,3-dimethylquinoxaline 1,4-dioxide (XXIXA) m. 249-50*. 5-Methoxy-2,3-dimethylquinoxaline 1,4-dioxide (XXIXA). m. 249-50*. 5-Methoxy-2,3-dimethylquinoxaline 1,4-dioxide (XXIXA)

the mixture filtered, and the filtrate acidified gave 5-hydroxy-2,3-dimethylquinoxaline 1-oxide, cream needles, m. 143-4.5° (from CRH6-cyclohexane); the reaction repeated in PhMO2 16 hrs. at 60-5°, the mixture cooled, treated with ice H2O and 10 cc. 10N NsOH, and the sparingly soluble Ns ealt filtered off and decomposed with dilute AcON gave

5-hydroxy-2.3-dimethylquinoxaline 1.4-dioxide, m. 171-3* (from CSH6). XXIXA (0.2 g.) in 0.5 cc. HNO3 (d. 1.4) and 0.5 cc. concentrated H2SO4, kept 1.5 hrs. at 0*, and the mixture poured on ice gave the 7(7)-nicro derivative, m. 250* (daccmposition) (from 50% AcOH). With 1:1 HNO3-H2O, XXIXA gave the HNO3 salt, m. 97* (deccmposition). XXIXA (1.42 g.), 50 cc. saturated M2HCO3, 5.25 g. iodine, and 70 cc. 10% KI kapt 10 days at room temperature, filtered, and the filtrate saturated with 502 gave 1.3 g. 7(7)-iodo derivative (XXX), m. 148-50*; a similar procedure gave the 7(7)-Br derivative, golden needles, m. 120* (deccmposition); XXXI loat iodine when recrystn. was attempted. XXIXA (2.06 g.) in 100 cc. saturated NAHCO3 and 3.7 g. Br in 30 cc. 15% KBr kept 0.25 hr. gave 1.26 g. di-Br derivative, dark red, m. 120* (explodes). XXVII (9 g.), 70 cc. 1.7M VIIIA containing 0.3% w/W H2SO4, and 0.1% NAH2O7 (XXXI) kept 7 hrs. at room temperature and 9 hrs. at 55*, concentrated in vacuo, and the residue treated with saturated NAHCO3 gave 4.5 g. 1.4-dioxide (XXXII) m. 134-5* (from CSH6); HCO3H and H2SO4 in Mc2CO were unsatisfactory for this oxidation while the addition of XXXI gave more consistent results than only VIIIA and H2SO4. XXXII and 10N NAOH kept 0.5 hr. at room temperature and acidified with HCl gave the 6-HO2CO analog, m. 243* (decomposition) (from H2CO4RCHOH). XXXII (2 g.) and 20 cc. McOH-NH3H kept 4 days at room temperature gave 0.55 g. 6-H2NCO analog, m. 266* (decomposition) (from H2CO4RCHOH). XXXII (2 g.) and 20 cc. McOH-NH3H kept 4 days at room temperature gave 0.55 g. 6-H2NCO compound-H2O, yellow needles, m. 215* (decomposition) (from M2CO4RCHOH). XXXII (6 cc. McOH-NH3H kept 4 days at room temperature gave 0.55 g. 6-H2NCO analog, m. 266* (decomposition) (from M2CO4RCHOH). XXII (7 cc. McOH-NH3H kept 10 hrs. at room temperature and acidified with HCl gave the 6-H02NCO analog-H2O, m. 215* (grom gave 10.5 cc. 2.1M VIIIA kept 10 hrs. at room temperature and 8 hrs. at 150* gave 2.7 1.4 dioxide (m. 150-22* (from H2CO4

107419-21-6 CAPLUS 6-Quinoxalinecarboxylic acid, 2,3-dimethyl-, 2-diethylaminoethyl ester (6CI) (CA INDEX NAME)

109939-89-1 CAPLUS 6-Quinoxalinecarboxamide, 2,3-dimethyl-N-(3-piperidinopropyl)-, 1,4-dioxide (6CI) (CA INDEX NAME)

(preparation of)
855619-47-3 CAPLUS
Isopentyl alcohol, {2,2'-biphenazine}-7,7'-dicarboxylate (5CI) (CA INDEX
NAME)

PAGE 1-B

- CH2- CH2- CHMe2

Ethyl alcohol, compd. with di-Et [2,2'-biphenazine]-7,7'-dicarboxylate (Sci) (CA INDEX NAME)

CM 1

CRN 858239-91-5 CMF C30 H22 N4 O4

CRN 64-17-5 CMF C2 H6 O

н3С-сн2-он

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.:

L13 ANSWER 179 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1955:8306 CAPLUS

AUTHOR(8): CORPORATE SOURCE: SOURCE:

43:1734e-i,1735a-b Newphenaxine derivatives and their tuberculostatic action Birkofer, Leonhard; Widmann, Arno Max-Planck-Inat., Heidelberg, Germany Chemische Berichte (1953), 86, 1295-1302 CODEN: CHBRAM; ISSN: 0009-2940 Journal

DOCUMENT TYPE:

L13 ANSWER 178 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
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05:40424 CAPLUS
ORIGINAL REFERENCE NO.: 50:7812N-1,7813a-d
TITLE:
Syntheses in the series of phenezine derivatives. I.
2,2'-Biphenezine and its derivatives

AUTHOR (S) : SOURCE:

Rozum, Yu. S.
Ukraine'kii Khemichnii Zhurnal (1955), 21, 491-5
CODEN: UKHZAS; ISSN: 0372-4190

boiled in 5-10% HCl, dried, and extracted with (CH2Cl)2, and the extract evaporated gave, after chromatographing twice in CSHS on Al2O3, 1.2 g. 2.2'-biphenazine (III), orange plates. m. 229* (from CSHS), insol. in REQ. H2O or petr. ether, giving brightly colored, readily hydrolyzed selts in concentrated mineral acids. Analogously was obtained: from I and c-O2NCGHAOMe, 5.5 9,9'-(MeO)2 derivative of III, red plates or prisms, m. 1a1* (from CSHS), dark red in acids, the color fading eventually; from 13,4-MeO (HANDCGH3)2 (IV) and II, 13,6* 4,4*-(HeO)2 derivative, dark red needles, m. 174* (from CSHS), red in acids; from c-CLCGHAO2 (V) and I and from V and IV the 9,9'-Cl2 (5.7%), red plates, m. 175* (from CSHS), red ben gave red acid solns. Similarly, I (36.8 g.) heated to 10° with 56 g. p-O2NCGHAME (VI) and 144 g. KOH, the product washed with ligroine and MeOH, the filter cake suspended in 3 1. H2O, heated to 0.5°, 86 g. Nenot added under egitation, the precipitated MnO2 removed by filtration, the filterate condensed to 300-400 mln, treated with C and glacial ACOH, and theven years.

sted, gave, after washing with EtOH and Et2O and drying, 9.5% 2,2"-biphenazine-7,7"-dicarboxylic acid, yellow needles, m. 320" (from glacial AcOH), soluble in bases and concentrated acids (yellow), insol.

common organic solvents; diamide (75%), m. 360-3° (decomposition) (fro HCONH2), insol. in common organic solvents, yellow-green in concentrat

HCONN2), insol. in common organic solvents, yellow-green in concentrated H2SO4; Me etcr (67%), orange plates. m. 210-12° (from SUOH and active C), insol. in EtOH and EtO, yellow in concentrated H2SO4; Et etcr (30%), red-orange needles, m. 130° (from EtOH), yellow in concentrated H2SO4; iso-amyl ester (34%), pink plates, m. 116° (from ino-AmoH), yellow in acid. Most of these retained 1-4 moles of crystallization solvent. Oth derivs. of III prepared were: 7,7°,4.7°M2(MeOI) (9.4%) (from IV, VI, and KOH), red plates, m. 238° (from MeCHCl2), blue in concentrated H2SO4; and 4,4°,9.9° (MeOI) (2.1%) (from IV, O-CANCSHAOMe, and KOH), red plates in column (310-390 m.) and H2SO4. Absolute maximum are given for all of these in tolumn (310-390 m.) and H2SO4 (410-610).

IT 855639-47-3, Isopentyl alcohol. (2,2°-biphemazine)-7,7°-dicarboxylate 88239-92-6. Ethyl alcohol, compound with di-Et [2,2°-biphemazine)-7,7°-dicarboxylate)

LANGUAGE: Unavailable

AB Some phenazine derivs, are prepared to be tested for their tuberculostatic action. Heating slowly 13 g. o-H2NC6H4CO2H, 16 g. o-O2NC6H4CO2H (1), and 25 g. finely powdered KOH to 65°, dissolving the melt in H2O, and concentrating the solution give dirk 1,6-phenazinedicerboxylate (11) (C.A. numbering), which on acidification gives 1.5 g. free acid, charring at 300-20° without melting (di-Et ester, prepared by dissolving 1.5 g. II in 20 cc. 100H H2SO4, pouring the solution into absolute EtoH, neutralizing the mixture with NaOH to pH 8-9, and extracting with ether, green-yellow needles.

300-20° without melting (di-Et ester, prepared by dissolving 1.5 g.
II in 20 cc. 100° H2504, pouring the solution into absolute ESCH, neutralizing the mixture with NaCH to pH 8-9, and extracting with ether, green-yellow needles,
m. 143°). Treating 0.6 g. Me 1-phenazinetoxylate with H2NOH
(from 2 g. HCl selt) gives 1-phenazinetoxylate with H2NOH
20°. Adding 3.5 g. 1-phenazinetoxoxoxylate with H2NOH
30°. NaOH and 2.5 g. Br and heating the mixture 5 min. at 70° give 744
1-aminophenazine, red needles, m. 176°, which (0.5 g.), refluxed
2.5 hrs. with 2 g. anhydrous glucose and 20 mg. NACH in 30 cc. absolute MaOH, gives 50° NrD-glucoside, vermilion needles, m. 185° (tetra-Ac derivative, prepared by heating 0.4 g. glucoside in 10 cc. CSHSN and 5 cc. Ac20
0.5 hr. on a water beth, orange needles, m. 181°). Reating 5 g. 1.
4.4 g. 2-CLOHYNH2, and 15 g. KOH to 80°, raising the temperature
5° every 20 min. until 130° is reached, keeping the mixture 3 hrs. at 130°, dissolving the cold melt in H2O, concentrating the filtered solution, and adding a little MeOH give the K benzo[a]phenazine-11-carboxylate (free acid, yellow needles, m. 256°). Mixing 2 g.
1,2-naphthoquinone (III) and 2.4 g. 3,4-(H2N) 2CSH2COZEt, each in 20 cc. AcOH, gives 80° Kt benzo[a]phenazine-0 for 10)-carboxylate (free acid, yellow needles, m. 36°. Macking
0.5 g. 1V and 6 cc. 40° N2H4.H2O in 15 cc. dioxane and 5 cc. EtoH 2 hrs. on a water bath gives the free acid, yellow needles, m. 36°. Heating
13,4-(H2N) 2CSH2CH2CH2CH2 in 10 cc. AcOH. gives 80° kg. H2NH2CSH2CH2CH2CH2 in 10 cc. AcOH. gives 80° kg. H2NH2CH2CH2CH2 in 1

L13 ANSWER 180 OF 181 CAPLUS COPYRIGHT 2006 ACS on STN
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ORIGINAL REFERENCE NO.: 48:11426-i,11427a-b

TITLE: AUTHOR(s): CORPORATE SOURCE: SOURCE:

Derivatives of 2-phenazinecarboxylic acid Pietra, Silvio; Maffei, Silvio; Rivolta, Angelamaria Univ. Pavia, Italy Annali di Chimica (Rome, Italy) (1953), 43, 227-31 CODBM. ANCRAI; ISSN: 0003-4592

Annali di Chimica (Rome, Italy) (1953), 43, 227-31 CODEN, ANCRAI; ISEN: 0003-4592

DOCUMENT TYPE: Journal LANGUAGE: Journal LANGUAGE: Journal LANGUAGE: Journal LANGUAGE: Journal LANGUAGE: Journal LANGUAGE: May a serie of 2-phenezinecarboxylic acid (I) has already been described [cf. ibid. 42, 519(1952)]. Various derive. are now described. Et2NH and MeOH with the acyl chloride of I yield 2-(diethylcarbamcyl)phenazine (II), m. 97.5-98*, and Me 2-phenazinecarboxylate (III) m. 153*.

NJH4 and III yield 2-phenazinecarboxylate (III) m. 153*.

NJH4 and III yield 2-phenazinecarboxylate acid hydrazide (IV), m. 269-70* (decomposition). IV and o-HoCOMe give actophenome are provided as a serie of the series of the seri

L13 ANSMER 181 OF 181

ACCESSION NUMBER:
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ACCESSION NUMBER:
OPERATE OF 181

A4:4690 CAPLUS

48:4690

A4:4690 Experimental psitacosis and lymphogranulome venereum (inguinale). II. The activity of guinoxaline 1,4-dioxide and substituted and related compounde, with a note on the morphological changes induced in lymphogranulome virus by these compounds and by antibiotics

AUTHOR(S):
BURGE:
BURGE:
CORPORATE SOURCE:
SOURCE:
Imperial Chem. Inde., Ltd., Manchester, UK
COSTORIAN OF 181 (1953), 8, 297-305

(1953), 8, 297-305 CODEN: BJPCAL; ISSN: 0366-0826

CODEN: BJPCAL; ISSN: 0366-0826

LANGUAGE: Journal

EAS Cf. C.A. 45, 3084h. Representative mono- and disubstituted quinoxaline

1,4-dioxides, substituted 2,3-dimethylquinoxaline 1,4-dioxides, phenarine
di-N-oxides, biquinoxaline tetra-N-oxides and N-oxides of miscellaneous

N-heterocyclic compds. possessed some degree of activity against the
largest viruses of the psittacosis-lymphogranuloma group. Quinoxaline
1,4-dioxide (1) and its substituted derivs. were most potent, the best

equaling Auracomycin in their activity against lymphogranuloma venerum in the mouse. Relatively few were active against this disease in the chick mathra and a second of the control of th

109939-89-1, 6-Quinoxalinecarboxamide, 2,3-dimethyl-N-(3-piperidinopropyl)-, 1,4-dioxide (effect on peittacosislymphogranuloma group viruses) 109939-89-1 CAPUE 6-Quinoxalinecarboxamide, 2,3-dimethyl-N-(3-piperidinopropyl)-, 1,4-dioxide (6CI) (CA INDEX NAME)

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